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## **PCT**

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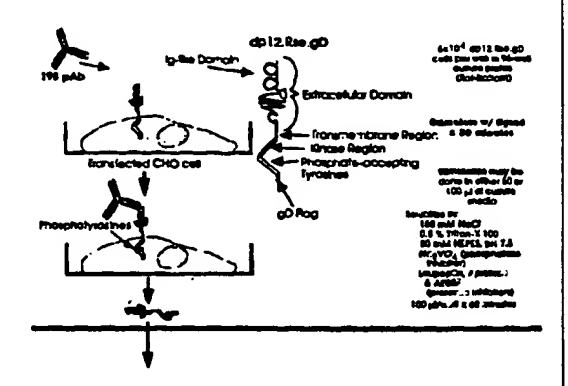
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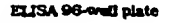
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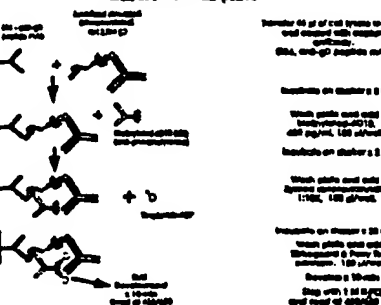
(54) Title: KINASE RECEPTOR ACTIVATION ASSAY

#### (57) Abstract

An assay for measuring activation (i.e., autophosphorylation) of a tyrosine kinase receptor of interest is disclosed. a) A first solid phase is coated with a substantially homogeneous population of cells so that the cells adhere to the first solid phase. The cells have either an endogenous tyrosine kinase receptor or have been transformed with DNA encoding a receptor or "receptor construct" and the DNA has been expressed so that the receptor or receptor construct is presented in the cell membranes of the cells. b) A ligand is then added to the solid phase having the adhering cells, such that the tyrosine kinase receptor is exposed to the ligand. c) Following exposure to the ligand, the adherent cells are solubilized, thereby releasing cell lysate. d) A second solid phase is coated with a capture agent which binds specifically to the tyrosine kinase receptor, or, in the case of a receptor construc, to the flag polypeptide. e) The cell lysate obtained in step c) is added to the wells containing the adhering capture agent so as to capture the receptor or receptor construct to the wells. f) A washing step is then carried out, so as to remove unbound cell lysate, leaving the captured receptor or receptor construct. g) The captured receptor or receptor construct is exposed to a labelled anti-phosphotyrosine antibody which identifies phosphorylated residues in the tyrosine kinase receptor. h) Binding of the anti-phosphotyrosine antibody to the captured receptor or receptor construct is measured.











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### KINASE RECEPTOR ACTIVATION ASSAY BACKGROUND OF THE INVENTION

#### Field of the Invention

This invention relates to a kinase receptor activation (KIRA) assay. 5 In particular, the invention relates to an assay for measuring autophosphorylation of the kinase domain of a receptor protein tyrosine (rPTK) using a kinase receptor activation, enzyme-linked kinase immunosorbent assay (KIRA ELISA).

#### Description of Related Art

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One mechanism for signal transduction in animals involves protein phosphorylation. Protein phosphorylation involves the action of protein kinase, an enzyme that transfers a phosphate group from a phosphate donor onto an acceptor amino acid in a substrate protein. Protein phosphatases provide a means for reversing the signal when the stimulus is removed.

Protein kinases have multiple substrates, and classification of the protein kinases is based on the acceptor amino acid specificity. The two most well characterized protein kinases are the protein kinases with a protein alcohol group as acceptor called protein serine/threonine kinases and the protein kinases with a protein phenolic group as acceptor called protein tyrosine kinases (Hunter, Methods in Enzymology 200:3-9 [1991]).

The most well known type of signal-transducing protein kinases are growth factor receptor protein tyrosine kinases (rPTKs). rPTKs usually comprise a large, glycosylated, extracellular ligand binding domain (ECD) and an intracellular domain (ICD) which contains a tyrosine kinase 25 catalytic domain. A single hydrophobic transmembrane (TM) domain connects the ECD and ICD. Examples of rPTKs include the insulin receptor, epidermal growth factor receptor (EGF-R), platelet-derived growth factor receptor (PDGF-R), insulin-like growth factor 1 receptor (IGF-1-R), and the HER2 receptor, to name a few. See, for example, Ullrich and Schlessinger Cell 30 61:203-212 (1990) and Fantl et al., Annu. Rev. Biochem. 62:453-481 (1993). rPTKs can phosphorylate exogenous protein substrates and intrinsic tyrosine residues via their catalytic tyrosine kinase domain. The intrinsic tyrosine residues normally reside in the ICD of the rPTK (see Figure 1 herein). Activation of the intracellular kinase domain of rPTKs appears to be mediated by receptor oligomerization which results from the conformational alteration of the ECD upon ligand binding thereto. See Ullrich and Schlessinger, supra.

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Serine-threonine kinases have also been disclosed in the literature. While most of the known protein serine-threonine kinases are cytoplasmic proteins, a family of mammalian transmembrane receptors with serine-5 threonine kinase domains has recently been found. Members of this receptor family have been described as binding TGF-B's and activin. For reviews of serine-threonine kinases, see Sale, G., Biochem. Soc. Transactions 20: 664-670 (1992); ten Dijke et al., Prog. in Growth Pactor Res. 5: 55-72 (1994); and Mathews, L., Endoc. Rev. 15(3): 310-325 (1994).

Various assays have been developed which measure tyrosine kinase activity. Some of these assays measure the ability of a tyrosine kinase enzyme to phosphorylate a synthetic substrate polypeptide. For example, an assay has been developed which measures growth factor-stimulated tyrosine kinase activity by measuring the ability of the kinase to catalyze the 15 transfer of the y-phosphate of ATP to a suitable acceptor substrate. See Pike, L., Methods of Enzymology 146:353-362 (1987) and Hunter, Journal of Biological Chemistry 257(9):4843-4848 (1982), for example. In this assay, the use of [y-32p]ATP permits the radioactive labeling of the phosphorylated substrate, which is a synthetic tyrosine-containing peptide. Others have described protein kinase assays wherein incorporation of 12p into a tyrosine kinase receptor, such as the EGF receptor (see Donato et al., Cell Growth Differ 3:259-268 [1992]), insulin receptor (see Kasuga et al., Journal of Biological Chemistry 257(17):9891-9884 [1982] and Kasuga et al., Methods in Enzymology 109:609-621 [1985]), and liver growth hormone receptor (see Wang et al., Journal of Biological Chemistry 267 (24):17390-17396 [1992]), is measured.

The discovery of anti-phosphotyrosine antibodies has provided a nonradioactive, alternative means for measuring phosphorylation of tyrosine residues. For example, White and Backer (Methods in Enzymology 201:65-67 [1991]) mention polyclonal antibodies which selectively bind to phosphotyrosine and are considered to be useful for studying rPTKs. An anti-phosphotyrosine monoclonal antibody was used in one of the assays referred to in Madden et al. (Anal Biochem 199:210-215 [1991]), which measured phosphatase activity toward the insulin receptor. Anti-35 phosphotyrosine antibodies were also used by Cleaveland et al., in their protein tyrosine kinase ELISA assay. See Cleaveland et al., Analytical Biochemistry 190:249-253 (1990). The method of Cleaveland et al. utilizes purified high-activity oncogene tyrosine kinases, v-src and v-fps, and measures the ability of these tyrosine kinases to phosphorylate synthetic polymeric substrates which are coated on an ELISA microtiter plate. The phosphotyrosine produced by src-induced ph sphorylation of the polymeric substrate is then quantitated by addition of an anti-phosphotyrosine antibody, the presence of which is detected using a second rabbit antimouse antibody which is linked to a reporter enzyme, horseradish peroxidase (HRPO). A similar ELISA assay has been developed by Lazaro et al., which is used for detection of a protein tyrosine kinase. See Lazaro et al., Analytical Biochemistry 192:257-261 (1991). Like the assay of Cleaveland et al., this assay also measures the ability of a protein tyrosine kinase to phosphorylate a synthetic substrate which is bound to microELISA wells.

A direct way to assess specific activation of rPTKs is by analysis of receptor autophosphorylation. See, e.g., Hunter and Cooper Ann Rev Biochem 54:897-930 (1985) and Ullrich and Schlessinger, Cell 61:203-212 (1990). Using this direct approach, Knutson and Buck disclose assays for measuring autophosphorylation of the insulin receptor under in situ or in vitro conditions (Archives of Biochemistry and Biophysics 285(2):197-204 [1991]). In the in situ assay, monolayer cultures of embryonic mouse 3T3-C2 fibroblasts (having the endogenous insulin receptor) are incubated with insulin in large cell culture dishes. Following incubation, the insulin receptor is extracted from the membranes. To achieve extraction of the insulin receptor, the cell monolayers are scraped into a buffer containing protease inhibitors and the cells are then disrupted in a homogenizer. The cellular homogenate is subsequently subjected to centrifugation for 60 min., and the pellet which forms is extracted into buffer containing Following a further centrifugation step, the supernatant (containing the insulin receptor) is incubated with an anti-insulin receptor antibody. Then, the receptor-antibody complex is incubated with protein A-agarose and unoccupied protein A sites are blocked with normal 30 rabbit IgG. The agarose beads are then centrifuged, the supernatants aspirated and the pellets are re-suspended in buffer containing the radiolabelled anti-phosphotyrosine antibody. The amount of bound iodinated anti-phosphotyrosine antibody is consequently measured.

Klein and his colleagues discuss an assay for measuring insulin activation of the insulin receptor (Klein et al., Diabetes 42:883-890 [1993]). In this assay, aliquots of a heterogeneous population of mononuclear blood cells (including T cells, B cells, macrophages etc) having the insulin receptor are exposed to insulin in centrifuge tubes.

The cells are then lysed in detergent using a motordriven homogenizer and the lysates are concentrated two- to fourfold using vacuum centrifugation. Sometimes, the insulin receptor is also partially purified using wheat germ agglutin agarose. The supernatants which form following centrifugation.

5 are then transferred to anti-insulin receptor-coated microtiter plates. Insulin (8.7nM) as well as kinase and phosphatase inhibitors are present during receptor immobilization in order to optimize the percentage of receptors captured to the microtiter plates. Activation of the insulin receptor is then measured by transphosphorylation of the substrate Poly
10 Glu, Tyr with 12P labeled ATP. The supernatants are then spotted onto absorbent paper and the paper is washed with cold TCA to remove unbound 12P
ATP. Remaining 12P-labeled Poly-Glu, Tyr on the washed absorbent paper is subsequently counted by scintillation counting.

in patients (Hagino et al., <u>Diabetes</u> 43:274-280 [1994]). As a first step in the assay, Hagino et al. stimulate a primary cell suspension, which is not particularly homogeneous with respect to cell type. In particular, heparinized blood (1ml washed twice with medium and resuspended in 1 ml of medium containing bovine serum albumin, BSA) is exposed to varying concentrations of insulin. The autophosphorylation reaction is stopped, the cells centrifuged for 30 min, the supernatant is discarded and the erythrocyte ghosts thus obtained are resuspended in buffer and centrifuged again. The pellet thereby obtained is adjusted to 500µl and solubilized in detergent. The solubilized materials are then centrifuged and the resulting supernatant is subjected to sandwich ELISA (using anti-insulin receptor antibodies to capture the insulin receptor) to determine the extent of insulin receptor autophosphorylation.

King et al. in Life Sciences 53: 1465-1472 (1993) describe a colorimetric assay for examining inhibitors of the epidermal growth factor 30 (EGF) receptor-associated tyrosine kinase in human intact epidermal A431 cells.

Several others have used an enzyme-conjugated form of the antiphosphotyrosine antibody in Western blot analyses which measure receptor
autophosphorylation. Briefly, Western blotting generally involves
electrophoresing activated rPTK on polyacrylamide gel. The rPTK is then
transferred to nitrocellulose and immunoblotted with the antiphosphotyrosine antibody which is labelled to enable detection. See, for
example, Wang, Molecular and Cellular Biology 5(12):3640-3643 (1985);

Glenney et al., Journal of Immunological Methods 109:277-285 (1988); Kamps, Methods in Enzymology 201:101-110 (1991); Kozma et al., Methods in Enzymology 201:28-43 (1991); Holmes et al., Science 256:1205-10 (1992); and Corfas et al., PNAS. USA 90:1624-1628 (1993). However, with Western blot analysis, accurate quantitation can be very tedious. Furthermore, this technique tends to be time- consuming and generally does not allow high sample throughput.

It is an object of the instant invention to provide a sensitive, reliable assay that measures receptor protein tyrosine kinase (rPTK) 10 autophosphorylation. The assay is desirably useful for qualitatively and quantitatively measuring kinase activation as well as facilitating identification and characterization of potential agonists and antagonists for a selected rPTK. It is a further object of the invention to provide an assay which enables ligand-receptor interactions to be studied for any selected rPTK.

This assay must have a capacity for high throughput, that is, the ability to reliably evaluate large numbers of samples in a relatively short period of time (e.g., in one day). The assay ideally does not use radioactive materials and is also amenable to automation.

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It is a further object, in at least one embodiment of the invention, to provide a generic assay which enables a rPTK of interest to be studied, regardless of whether or not a receptor-specific capture agent having the desired characteristics is available. Furthermore, it is an object of the invention to provide an assay which substantially represents the activity 25 of the tyrosine kinase receptor in situ. This is desirable insofar as it reduces the possibility that altered interactions between the receptor and the ligand may occur as a consequence of the receptor not being membranebound. Furthermore, if the receptor is a multimeric complex, this assay enables the correctly assembled receptor to be studied. additional object to provide a method for measuring serine-threonine kinase phosphorylation, phosphorylation of intracellular kinases and phosphatase activity.

These and other objects will be apparent to the ordinary artisan upon consideration of the specification as a whole.

#### SUMMARY OF THE INVENTION

Accordingly, the invention provides an assay for measuring activation (i.e., autophosphorylation) of a tyrosine kinase receptor of interest.

The assay can be divided into two major stages, each of which is generally performed in separate assay plates. The first stage of the assay involves activating the receptor and is termed the kinase receptor activation (KIRA) stage of the assay. The second stage of the assay involves measuring receptor activation. Conveniently, this is achieved using an enzyme-linked immunosorbent assay (ELISA) to measure receptor activation.

The KIRA stage of the assay involves activating a tyrosine kinase receptor which is located in the cell membrane of an eukaryotic cell such that the extracellular domain of the receptor faces the external milieu of the cell, the transmembrane domain is located in the cell membrane and the kinase domain is located intracellularly. This stage of the overall assay involves steps (a) to (c) below:

(a) The first solid phase (e.g., a well of a first assay plate) is coated with a substantially homogeneous population of cells (usually a mammalian cell line) so that the cells adhere to the solid phase. Often, the cells are adherent and thereby adhere naturally to the first solid phase. In one embodiment of the invention, the cells have an endogenous tyrosine kinase receptor presented in the cell membrane as discussed above.
20 In an alternative embodiment, the cells have been transformed with DNA encoding a tyrosine kinase receptor or a "receptor construct" defined further below, which DNA is expressed by the cells such that the receptor or receptor construct is suitably positioned in the cell membranes thereof.

The receptor construct comprises a fusion of a kinase receptor and a flag polypeptide. The flag polypeptide is recognized by the capture agent, often a capture antibody, in the ELISA part of the assay. Use of a receptor construct as disclosed herein is particularly advantageous since it provides a "generic" assay wherein autophosphorylation of any tyrosine kinase receptor can be measured, regardless of whether or not a receptor-specific capture agent having the required characteristics is available. Often, the tyrosine kinase receptor is a fusion protein comprising the ECD of a selected tyrosine kinase and the catalytic ICD (and possibly the transmembrane domain) of another well characterized tyrosine kinase (e.g., the Rse receptor).

(b) An analyte is then added to the wells having the adhering cells, such that the tyrosine kinase receptor is exposed to (or contacted with) the analyte. This assay enables identification of agonist and antagonist ligands for the tyrosine kinase receptor of interest. In order to detect

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the presence of an antagonist ligand which blocks binding and/or activation of the receptor by an agonist ligand, the adhering cells are exposed to the suspected antagonist ligand first and then to the agonist ligand (or to a mixture of the agonist and antagonist) so that competitive inhibition of receptor binding and activation can be measured. Also, the assay can identify an antagonist which binds to the agonist ligand and thereby reduces or eliminates its ability to bind to, and activate, the rPTK. To detect such an antagonist, the suspected antagonist and the agonist for the rPTK are incubated together and the adhering cells are then exposed to this mixture of ligands.

(c) Following exposure to the analyte, the adhering cells are solubilized using a lysis buffer (which has a solubilizing detergent therein) and gentle agitation, thereby releasing cell lysate which can be subjected to the ELISA part of the assay directly, without the need for 15 concentration or clarification of the cell lysate. Thus, this assay provides a significant improvement over assays described by Knutson and Buck, supra, Klein et al., supra, and Hagino et al., supra, insofar as it is surprisingly unnecessary to concentrate the cell lysate prior to the ELISA. Furthermore, unlike the other assays, in the instant assay the cells can be lysed in lysis buffer using gentle agitation without the need for homogenizing, centrifuging or clarifying the cells. The cell lysate thus prepared is then ready to be subjected to the ELISA stage of the assay. It has been discovered that, surprisingly, the first assay plate can be stored at freezing temperatures (i.e., at about -20° to -70°C) for significant periods of time (at least 6 months) before the ELISA stage of the assay. This is a significant finding insofar as the KIRA and ELISA stages of the assay can be performed on separate days.

The ELISA component of the assay comprises steps (d) to (h), described below.

(d) As a first step, the second solid phase (usually a well of an ELISA microtiter plate) is coated with a capture agent (often a capture antibody) which binds specifically to the tyrosine kinase receptor, or, in the case of a receptor construct, to the flag polypeptide. Coating of the second solid phase is carried out so that the capture agent adheres to the second solid phase. The capture agent is generally a monoclonal antibody, but, as is described in the examples herein, polyclonal antibodies may also be used.

- (e) The cell lysate obtained in step (c) of the above-mentioned KIRA stage of the assay is exposed to, or contacted with, the adhering capture agent so that the receptor or receptor construct adheres to (or is captured in) the second solid phase. Unlike the assay of Klein et al., the instant 5 assay does not require the ligand for the receptor as well as kinase inhibitors to be present to achieve suitable immobilization of the receptor or receptor construct to the second solid phase.
  - (f) A washing step is then carried out, so as to remove unbound cell lysate, leaving the captured receptor or receptor construct.

- (g) The adhering or captured receptor or receptor construct is then exposed to, or contacted with, an anti-phosphotyrosine antibody which identifies phosphorylated tyrosine residues in the tyrosine kinase receptor. In the preferred embodiment, the anti-phosphotyrosine antibody is conjugated (directly or indirectly) to an enzyme which catalyses a color 15 change of a non-radioactive color reagent. Accordingly, phosphorylation of the receptor can be measured by a subsequent color change of the reagent. The enzyme can be bound to the anti-phosphotyrosine antibody directly, or a conjugating molecule (e.g., biotin) can be conjugated to the antiphosphotyrosine antibody and the enzyme can be subsequently bound to the 20 anti-phosphotyrosine antibody via the conjugating molecule.
  - (h) Finally, binding of the anti-phosphotyrosine antibody to the captured receptor or receptor construct is measured, e.g., by a color change in the color reagent.

The invention also pertains to a Rse.flag reagent which is 25 particularly useful for use in the KIRA ELISA assay. The Rse.flag reagent is a polypeptide comprising a fusion of a flag polypeptide (usually the gD flag described herein) to the carboxyl terminus of the intracellular domain of the Rse rPTK. Generally, the transmembrane domain of Rse and the extracellular domain of another rPTK of interest are also present in the fusion polypeptide reagent. The nucleic acid encoding this reagent and a cell transformed therewith are also claimed.

In yet a further aspect, the invention relates to a kit which can be used in the KIRA ELISA disclosed above which comprises an anti-flag polypeptide capture agent (e.g. a capture antibody) which is usually bound 35 to the second solid phase as described herein. Thus, the kit generally provides an ELISA microtiter plate having an anti-flag polypeptide capture antibody adhering to a well thereof. Optionally, the kit also provides an anti-phosphotyrosine antibody which is often labelled, or reagents for labelling the anti-phosphotyrosine antibody are supplied with the kit. Sometimes, a homogeneous population of cells which have been transformed with a receptor construct as described herein are also provided with the kit. The kit also suitably includes instructions for carrying out the KIRA ELISA.

#### BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1A-1C are diagrammatic representations of Rse.gD (Figure 1A),
Receptor ECD/Rse.gD chimera (Figure 1B) and a CHO cell transformed with the
Receptor ECD/Rse.gD chimera (Figure 1C).

Figures 2A and 2B depict an alignment of the amino acid sequence (SEQ ID NO: 1) and nucleotide sequence (SEQ ID NO: 2) of Rse.gD. The residues of the signal sequence are indicated with an (\*), the transmembrane domain of Rse is boxed and the ECD and ICD of Rse are also delineated. The residues of the gD flag sequence are underlined.

Figure 3 is a flow diagram of an exemplary strategy for selecting a suitable capture agent for use in the assay.

Figure 4 is a flow diagram of an exemplary strategy for selecting a transformed cell suitable for use in the assay, where the cell has a receptor construct with an amino-terminal flag polypeptide located in the cell membrane thereof.

Figure 5 is a flow diagram of an exemplary strategy for selecting a transformed cell suitable for use in the assay, where the cell has a receptor construct with a carboxyl-terminal flag polypeptide located in the cell membrane thereof.

Figure 6 is a flow chart and cartoon illustrating the KIRA ELISA assay for the HER2 receptor described in Example 1.

Figure 7 depicts a pl85<sup>HER2</sup>/HRGβl<sub>177-244</sub> KIRA ELISA standard curve obtained using the assay described in Example 1. To obtain the standard curve, MCF-7 cells (2x10<sup>5</sup>) were stimulated with 3000, 1000, 333, 111, 37, 12, 4, or 0 pM HRGβl<sub>177-244</sub>, as determined by quantitative amino acid analysis (q.a.a.a.). Each calibrator concentration was run in triplicate. The values derived from 10 such standard curves were averaged (total n = 30) and are presented as mean ABS<sub>450/650</sub> ± sd vs. HRGβl<sub>177-244</sub> concentration.

Figure 8 depicts heregulin specificity of p185<sup>HER2</sup>/HRG KIRA ELISA of Example 1. In the assay, MCF-7 cells (2x10<sup>5</sup>) were stimulated with either HRGβ1<sub>177-244</sub> (■) at 3000, 1000, 333, 111, 37, 12, 4 or 0 pM or IGF-1(•), EGF (□), VEGF (●) or insulin (♦) at 30000, 10000, 3333, 1111, 370, 120, 40 or

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0 pM. For all concentrations of ligands, n = 3 and data are presented as average ABS<sub>450/650</sub> ± sd vs. ligand concentration.

Figure 9 is a flow chart and cartoon illustrating the KIRA ELISA assay for the Rse receptor described in Example 2.

Figure 10 depicts a Rse KIRA ELISA standard curve obtained using the assay described in Example 2. To obtain the standard curve, CHO cells transformed with the Rse.gD construct were stimulated with 1:100, 1:200, 1:400, 1:800, 1:1600, 1:3200 or 0 diluted, anti-Rse agonist antibody. Each calibrator concentration was run in triplicate. The values are presented 10 as mean ABS<sub>450/650</sub> ± sd vs. 1/dilution agonist antibody.

Figure 11 is a flow chart and cartoon illustrating the KIRA ELISA assay for the trk receptors (i.e., trk A, trk B, and trk C) described in Example 3.

Pigures 12A-12D depict an alignment of the amino acid acid sequence (SEQ ID NO: 3) and nucleotide sequence (SEQ ID NO: 4) of gD.trk A used in the assay described in Example 3. The residues of the signal sequence are indicated with an (\*), the residues of the gD flag sequence are underlined, the residues of the transmembrane domain of trk A are in bold and the ECD and ICD thereof are also delineated.

Figures 13A-13D depict an alignment of the amino acid sequence (SEQ ID NO: 5) and nucleotide sequence (SEQ ID NO: 6) of gD.trk B used in the assay described in Example 3. The residues of the signal sequence are indicated with an (\*), the residues of the gD flag sequence are underlined, the residues of the transmembrane domain of trk B are in bold and the ECD and ICD thereof are also delineated.

Figures 14A-14D depict an alignment of the amino acid sequence (SEQ ID NO: 7) and nucleotide sequence (SEQ ID NO: 8) of gD.trk C used in the assay described in Example 3. The residues of the signal sequence are indicated with an (\*), the residues of the gD flag sequence are underlined, the residues of the transmembrane domain of trk C are in bold and the ECD and ICD thereof are also delineated.

Figures 15A-15C depict standard curves for trk A, B and C, respectively, which were obtained using the assay described in Example 3. To obtain the standard curves, CHO cells transformed with the gD.trk constructs were stimulated with 3000, 1000, 333, 111, 37, 12, 4 or 0 pM of ligand, i.e. nerve growth factor (NGF, □), neurotrophin 3 (NT3, •) or neurotrophin 5 (NT5, .). The values are presented as mean ABS450/650 ± sd vs. ligand concentration.

Figures 16A-16L depict the nucleotide sequence (SEQ ID NO: 9) of the pSVI17.ID.LL expression vector used for expression of Rse.gD in Example 2.

Figure 17 is a diagrammatic representation of the MPL/Rse.gD chimeric receptor described in Example 4.

Figure 18 is a flow chart and cartoon illustrating the KIRA ELISA for the MPL/Rse.gD chimeric receptor described in Example 4.

#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

#### Abbreviations and Definitions

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"rPTK" means a receptor protein tyrosine kinase.

"ECD", "TM domain" and "ICD" refer to the extracellular domain, transmembrane domain and intracellular domain of a rPTK, respectively.

"Kinase Receptor Activation" or "KIRA" when used throughout this application refers to the first stage of the instantly claimed assay wherein a cell-bound rPTK is exposed to a potential agonist/antagonist ligand which may (or may not) induce phosphorylation of tyrosine residues in the intracellular domain of the rPTK. The KIRA is generally carried out in the "first assay plate" as defined herein.

"Enzyme-Linked Immunosorbent Assay" or "ELISA" refers to the second stage of the instantly claimed assay and involves measuring tyrosine phosphorylation of the rPTK. The ELISA is normally carried out in the "second assay plate" as disclosed in this application. The ELISA is a "sandwich ELISA" insofar as it involves capturing the rPTK or receptor construct to the second solid phase (usually the well of an ELISA microtiter plate). ELISA assays generally involve the preparation of enzyme-antibody conjugates. The conjugated enzyme cleaves a substrate to generate a colored reaction product that can spectrophotometrically. In this assay, the absorbance of the colored solution in individual microtiter wells is proportional to the amount of phosphotyrosines. A review of ELISA is found in <u>Current Protocols in</u> 30 Molecular Biology, Vol. 2, chapter 11 (1991). While the term "ELISA" is used to describe the second stage of the instant assay, it is only a preferred embodiment of the invention, since, as disclosed herein, techniques other than enzymatic detection are available for measuring binding of the anti-phosphotyrosine antibody to the activated receptor.

The terms "receptor", "kinase receptor", "tyrosine kinase", "tyrosine kinase receptor", "receptor protein tyrosine kinase" and "rPTK" are used

interchangeably herein and refer to a protein having at least one phosphate accepting phenolic group. The protein is usually a receptor insofar as it has a ligand-binding ECD, TM domain and ICD. The ICD usually comprises a catalytic kinase domain and has one or more phosphate accepting tyrosine 5 residues. See Figures 1A and 1B, for example. Examples of tyrosine kinase receptors include the insulin receptor, insulin related receptor, epidermal growth factor receptor (EGF-R), platelet-derived growth factor receptors A and B (PDGF-R-A and PDGF-R-B). insulin-like growth factor 1 receptor (IGF-1-R), macrophage colony-stimulating factor receptor (M-CSF-R), 10 HER2/neu/c-erbB-2 receptor, HER3/c-erbB-3 receptor, Xmrk receptor, IRR receptor, fibroblast growth factor (FGF) receptors bek and flg, c-kit receptor, Flk/kDR receptor, Rse receptor, the Eph, Elk, Eck, Eek, Erk, Cek4/Mek4/HEK and Cek5 receptors, Axl receptor, hepatocyte growth factor receptor (HGF-R), Flt1 VEGF receptor, SAL-S1 receptor, HpTK 5 receptor, 15 trkA receptor, trkB receptor, and trkC receptor. See, for example, Ullrich and Schlessinger Cell 81:203-212 (1990); Fantl et al., Annu. Rev. Biochem. 62:453-481 (1993); Mark et al., Journal of Biological Chemistry 269(14):10720-10728 (1994); and WO 93/15201.

The terms mentioned above encompass chimeric "receptor" molecules which comprise at least the extracellular domain of a selected tyrosine kinase and the intracellular domain, and optionally, the transmembrane domain of another tyrosine kinase. Of course, the tyrosine kinase of interest can provide the transmembrane domain and/or intracellular domain. The terms also encompass amino acid sequence variants and covalent derivatives of the various rPTKs provided they still display tyrosine kinase phosphorylation activity in the KIRA ELISA. Therefore, the variants will general have conservative amino acid alterations. The individual domains of the tyrosine kinase can be delineated based on sequence homology to known tyrosine kinases and hydrophobicity plots. For example, the hydrophobic transmembrane domain can be readily determined and the ECD and ICD are usually amino-terminal and carboxyl terminal to the transmembrane domain, respectively. Conveniently, the transmembrane domain and ICD of the Rse receptor can be fused to the ECD of a tyrosine kinase of interest, thereby forming a chimeric receptor which is encompassed by the terms denoting a receptor as mentioned above.

In the preferred embodiment, the rPTK is selected from the group consisting of HER2 receptor (Ullrich and Schlessinger, supra), Rse receptor

(Mark et al., supra and SEQ ID NO: 1), trk A recept r (SEQ ID NO: 3), trk B receptor (SEQ ID NO: 5) and trk C receptor (SEQ ID NO: 7).

By "autophosphorylation" is meant activation of the catalytic kinase domain of the rPTK, whereby at least one intrinsic tyrosine residue is phosphorylated. Generally, autophosphorylation will result when an agonist molecule binds to the extracellular domain of the kinase receptor. Without being limited to any particular mechanism of action, it is thought that binding of the agonist molecule may result in oligomerization of the kinase receptor which causes activation of the catalytic kinase domain.

By "solid phase" is meant a non-aqueous matrix to which the cells (in the KIRA stage of the assay) or the capture agent (in the ELISA stage of the assay) can adhere. Usually, the solid phase comprises the well of an assay plate but the invention is by no means limited to this embodiment. For example, the solid phase can comprise a discontinuous solid phase of discrete particles. The particles can be porous and formed from a number different materials, e.g., polysaccharides (e.g. agarose), polyacrylamides, polystyrene, polyvinyl alcohol, silicones and glasses. For examples of suitable particulate solid phases, see U.S. Patent No. 4,275,149.

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By "well" is meant a recess or holding space in which an aqueous sample can be placed. The well is provided in an "assay plate". The invention usually employs a "first assay plate" which is formed from a material (e.g. polystyrene) which optimizes adherence of cells (having the receptor or receptor construct) thereto. Generally, the individual wells 25 of the first assay plate will have a high surface area to volume ratio and therefore a suitable shape is a flat bottom well (where the cells are adherent). The "second assay plate" is generally formed from a material (e.g. polystyrene) which optimizes adherence of the capture agent thereto. The second assay plate may have the same general construction and/or characteristics as the first assay plate. However, separate plates are used for the KIRA stage of the assay and the ELISA stage of the assay.

In the preferred embodiment of the invention, both the first assay plate and the second assay plate are "microtiter" plates. "microtiter" plate when used herein refers to an

35 assay plate having between about 30 to 200 individual wells, usually 96 wells. Often, the individual wells of the microtiter plate will hold a maximum volume of about 250 $\mu$ l. Conveniently, the first assay plate is a 96 well polystyrene or plastic, cell culture microtiter plate (such as that

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sold by Becton Dickinson Labware, Lincoln Park, NJ), which allows for automation. Often, about  $50\mu$ l to  $300\mu$ l, more preferably  $100\mu$ l to  $200\mu$ l, of an aqueous sample comprising cell culture media with the cells suspended therein will be added to each well of the first assay plate in the KIRA stage of the assay. It is desirable to seed between about 1 x 104 to 3 x 105 cells per well. More preferably, 5 x 104 to 1 x 105 cells per well are Usually, the second assay plate will comprise a polystyrene microtiter ELISA plate such as that sold by Nunc Maxisorp, Inter Med, Denmark.

The term "homogeneous population of cells" refers to a substantially homogeneous population of cells wherein at least about 80%, and preferably about 90%, of the cells in the population are of the same cell type. Therefore, it is convenient to use a cell line. The cell line is a eukaryotic cell line, normally an animal cell line and desirably a 15 mammalian cell line.

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The cells have, or are transformed to produce, the selected receptor or a receptor construct. For example, where the kinase receptor is known to be present in a certain cell line (e.g., the HER2 receptor in the MCF-7 cell line) no transformation step is required. Conversely, it may be 20 necessary to transform a cell with a nucleic acid encoding the receptor, where the cell does not make the receptor, or does not have suitable numbers of the receptor in the cell membrane thereof. Accordingly, the cell is transformed with a nucleic acid encoding the receptor (or receptor construct) and the nucleic acid is expressed so that the ECD of the receptor faces the external milieu of the cell, the transmembrane domain is located in the cell membrane and the kinase domain is located intracellularly.

Where the assay relies on activating the endogenous rPTK, a cell line is selected which is known to produce the rPTK of interest, provided 30 sufficient levels of the rPTK are present in the cell membrane thereof to enable detection. As a general proposition, a minimum number of about 1 x 104 receptors/cell is required. For example, the MCF-7 cell line (ATCC-HTB 22) which produces the HER2 receptor was shown to be useful in the assay. There are 5 x 104 HER2 receptors/MCF-7 cell. Examples of other cell lines 35 and their respective rPTKs include, embryonic mouse 3T3-C2 fibroblast cell line and the insulin receptor, and Hep 3B (ATCC # HB 8064) cell line and the Rse receptor. However, the degree of expression of the rPTK nucleic acid in the cell line is not so high that it results in constitutive

phosphorylation of the rPTK. For example, the SK-BR-3 cell line (ATCC HTB30), which has  $3 \times 10^6$  HER2 receptors/cell, was found to be unsuitable for use in the assay disclosed herein. Therefore, it may be useful to use a cell line which has less than about 3 x 106 receptors/cell, depending on 5 the type of receptor. The number of receptors/cell can be measured using Scatchard analysis, for example (Scatchard, Ann. NY Acad. Sci. 51:660-672 [1949]; and Goodwin et al., Cell 73:447-456 [1993]). However, selection of a cell line having a suitable number of receptors/cell is possible using the techniques described herein.

The term "adherent" when used herein to describe the cell, refers to a cell which naturally adheres to the first solid phase (often the well of the first assay plate), thereby forming a fairly uniform coating of the cells on the inside surface of the well. The uniform coating of cells generally forms following incubation of the cells in the wells of the first assay plate for about 8-16 hours. After incubation, non-adhering cells and cell culture medium are decanted off the first assay plate. Incubation is usually carried out at a temperature which is optimal for cell growth, i.e. about 37°C. Examples of adherent cell lines include CHO cells (Urlaub and Chasin, Proc. Natl. Acad. Sci. USA 77:4216 [1980]), MCF-7 cells (ATCC HB 20 22), 293 cells (Graham et al., J. Gen Virol. 36:59 [1977]), Swiss albino 3T3 fibroblast cell line (ATCC No. CCL 92) and U937 macrophage cell line (ATCC No. CRL 1593).

A "flag polypeptide" comprises a short polypeptide which has enough residues to provide an epitope (preferably a linear epitope) against which a "capture agent" thereagainst can be made, yet is short enough such that it does not interfere with activity of the rPTK. The flag polypeptide is also sufficiently unique so that the capture agent thereagainst does not bind to other reagents in the assay. Selection of a "unique" flag polypeptide sequence can be accomplished by comparing the sequence of a proposed flag polypeptide against other known sequences in Genbank or EMBL, for example. Suitable flag polypeptides generally have at least 6 amino acid residues and usually between about 8-80 amino acid residues (preferably between about 9-30 amino acid residues).

By "receptor construct" is meant a polypeptide which comprises a fusion of a kinase receptor and a flag polypeptide as defined above. The flag polypeptide is provided at a location in the receptor construct such that: a) the flag polypeptide does not interfere with ligand binding to the receptor; b) the flag polypeptide does not interfere with

autophosphorylation of the receptor and c) the flag polypeptide is presented in a suitable configuration so that it can bind to the capture agent in the ELISA stage of the assay. Often, the polypeptide flag will be present at the N-terminus f the receptor construct. See, for example, Example 3 which refers to the gD.trk constructs. Alternatively, the flag polypeptide may be present at the C-terminus of the receptor construct. See, for example, Example 2 which refers to the Rse.gD construct. See also Figures 1A-1C. The Rse construct disclosed herein is particularly useful, since the ICD (and optionally the transmembrane domain) thereof can be fused to the ECD of a kinase receptor of interest, thereby obviating the need to establish where the flag polypeptide should be located with respect to the kinase receptor of interest.

"Rse.gD" refers to a receptor construct which is the Rse receptor protein tyrosine kinase with the Herpes Simplex virus glycoprotein D (gD) 15 flag polypeptide fused to the COOH-terminus thereof.

"Rse.flag reagent" refers to a polypeptide which comprises the ICD of the Rse receptor fused at its COOH-terminus to a flag polypeptide (normally the gD flag polypeptide). Sometimes, the TM domain of Rse and the ECD of a rPTK of interest will also be present in the Rse.gD. reagent. "Receptor ECD/Rse.gD Chimera" refers to a fusion of the ECD of a rPTK of interest to the TM and ICD domains of Rse which are fused COOH-terminally to the gD flag polypeptide.

"gD.trkA", "gD.trkB" and "gD.trkC" refer to each of the trk receptors (A-C) having the gD flag polypeptide fused to the amino-termini thereof.

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By "capture agent" is meant a compound or agent which is able to adhere to the second solid phase, as herein defined, and which is selective for a rPTK or receptor construct. Thus, the capture agent captures the receptor or receptor construct to the wells of the second assay plate. Usually, the capture agent binds selectively to the flag polypeptide which 30 has been fused to the receptor of interest. Binding of the capture agent is not affected by the presence or absence of ligand bound to the receptor and does not induce receptor activation upon capture. Furthermore, the capture agent does not sterically block access to the phosphorylated tyrosine(s) by the anti-phosphotyrosine antibody. Means for selecting 35 suitable capture agents are described herein. Generally, the capture agent will comprise an antibody (e.g., an affinity purified polyclonal antibody or a monoclonal antibody), but other selective agents, such as streptavidin which binds selectively to the "strep-tag" polypeptide can also be used

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Schmidt et al., Protein Engineering 6(1):109-122 [1993]). Streptavidin can be purchased commercially from Zymed Laboratories, S. San Francisco, CA, for example. Alternatively, the capture agent can comprise protein A (which binds specifically to immunoglobulins). In this embodiment of the invention, the activated receptor or receptor-construct present in the cell lysate is incubated with an antibody which binds specifically thereto, thereby forming a receptor-antibody complex. This complex can be captured by protein A by virtue of its specific binding to the antibody present in the complex. Protein A can be purchased commercially from Pharmacia Biotech, Inc., Piscataway, New Jersey, for example. A strategy for selecting a suitable capture agent is depicted in Figure 3 and will be described in more detail later herein.

In the most preferred embodiment, the capture agent is a monoclonal antibody which binds specifically to a flag polypeptide (which is present in the receptor construct). Examples of suitable flag polypeptides and their respective capture antibodies include the flu HA flag and its antibody 12CA5, (Field et al., Mol. Cell. Biol. 8:2159-2165 [1988]); the c-myc flag and the 8F9, 3C7, 6E10, G4, B7 and 9E10 antibodies thereto (Evan et al., Molecular and Cellular Biology 5(12):3610-3616 [1985]); as well as 20 the Herpes Simplex virus glycoprotein D (gD) flag and the 5B6 antibody thereto (Paborsky et al., Protein Engineering 3(6):547-553 [1990] and Mark et al., Journal of Biological Chemistry 269 (14):10720-10728 [1994]). Other flag polypeptides have been disclosed. Examples include the Flag-peptide (Hopp et al., BioTechnology 6:1204-1210 [1988]); the KT3 epitope peptide (Martin et al., Science 255:192-194 [1992]); an  $\alpha$ -tubulin epitope peptide (Skinner et al., J. Biol. Chem 266:15163-15166 [1991]); and the T7 gene 10 protein peptide tag (Lutz-Freyermuth et al., Proc. Natl. Acad. Sci. USA 87:6393-6397 [1990]). Once the flag polypeptide has been selected as discussed above, a capture antibody thereto can be generated using the 30 techniques disclosed herein.

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The term "analyte" refers to a compound or composition to be studied, usually to investigate its ability to activate (or prevent activation of) the tyrosine kinase receptor of interest. The analyte can comprise a bodily fluid (such as plasma or amniotic fluid) or a composition known to 35 contain, or suspected of containing, a ligand for the tyrosine kinase receptor. The analyte can also comprise a cell which has a ligand to the rPTK of interest.

"Ligand" when used herein refers to a molecule which is able to bind to the ECD of the tyrosine kinase of interest or to a known agonist for the tyrosine kinase of interest. The ligand will usually be an agonist or antagonist f r the tyrosine kinase.

By "agonist" is meant a molecule which is able activate the intracellular kinase domain of the tyrosine kinase upon binding to the ECD. Often, the agonist will comprise a growth factor (i.e., a polypeptide that is able to stimulate cell division). Exemplary growth factors include heregulin (HRG), insulin, insulin-like growth factors I and II (IGF-I and IGF-II), epidermal growth factor (EGF), interleukins (e.g., IL-8), macrophage colony-stimulating factor (M-CSF), erythropoietin (EPO), platelet-derived growth factor (FDGF), fibroblast growth factor (FGF), transforming growth factors alpha and beta (TGF-α and TGF-β), hepatocyte growth factor (HGF), and nerve growth factor (NGF). Alt-4rnatively, the agonist can be an antibody against the rPTK (see, e.g., Yarden, Proc. Natl. Acad. Sci., USA 87:2569-2573 [1990]). However, other non-protein agonists such as small organic molecules are also encompassed by the invention.

Usually, the antagonist will either: (a) bind to the rPTK and thereby block binding and/or activation of the rPTK by an agonist thereto (the antagonist may bind to the ECD of the rPTK, but this is not necessarily the case) or (b) bind to the agonist and thus prevent activation of the rPTK by the agonist. This assay facilitates the detection of both types of antagonist. The antagonist may, for example, comprise a peptide fragment comprising the receptor binding domain of the endogenous agonist ligand for the receptor. The antagonist may also be an antibody which is directed against the ECD of the rPTK, or against a known agonist for the rPTK. However, other non-protein molecules are also encompassed by this term.

The term "antibody" is used in the broadest sense and specifically covers monoclonal antibodies and antibody compositions with polyepitopic specificity (i.e. polyclonal antibodies). The polyclonal antibodies are preferably "affinity purified" antibodies. The term "affinity purified" means that the antibodies have been purified using the antigen (e.g. the rPTK or fragment thereof or the flag polypeptide) to selectively purify the polyclonal antibodies. Affinity purification can be achieved by immobilizing the antigen on an affinity column (e.g. an agarose column) and passing the polyclonal antibodies through the column. The affinity purified antibodies can be subsequently eluted from the column by changing

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the elution conditions or by adding a chaotropic agent, for example. For a review of affinity purification techniques with respect to antibodies, see Current Protocols in Immunology, Ed. Coligen et al., Wiley publishers, Vols. 1 and 2, for example.

The term "monoclonal antibody" as used herein refers to an antibody obtained from a population of substantially homogeneous antibodies. 1.e., the individual antibodies comprising the population are identical except for possible naturally-occurring mutations that may be present in minor amounts. Monoclonal antibodies are highly specific, being directed against a single antigenic site. Furthermore, in contrast to conventional (polyclonal) antibody preparations which typically include different antibodies directed against different determinants (epitopes), each monoclonal antibody is directed against a single determinant on the antigen.

The monoclonal antibodies herein include hybrid and recombinant antibodies produced by splicing a variable (including hypervariable) domain of a selected antibody with a constant domain (e.g. "humanized" antibodies), or a light chain with a heavy chain, or a chain from one species with a chain from another species, or fusions with heterologous 20 proteins, regardless of species of origin or immunoglobulin class or subclass designation, as well as antibody fragments (e.g., Fab, F(ab')2, and Fv), so long as they exhibit the desired biological activity. [See, e.g. U.S. Patent No. 4,816,567 and Mage & Lamoyi, in Monoclonal Antibody Production Techniques and Applications, pp.79-97 (Marcel Dekker, Inc., New York (1987)].

Thus, the modifier "monoclonal" indicates the character of the antibody as being obtained from a substantially homogeneous population of antibodies, and is not to be construed as requiring production of the antibody by any particular method. For example, the monoclonal antibodies to be used in accordance with the present invention may be made by the hybridoma method first described by Kohler & Milstein, Nature, 256: 495 (1975), or may be made by recombinant DNA methods (U.S. Patent No. 4,816,567). The "monoclonal antibodies" may can also be isolated from phage antibody libraries using the techniques described in Clackson et al., 35 Nature, 352: 624-628 (1991) and Marks et al., J. Mol. Biol., 222: 581-597 (1991), for example.

The term "anti-phosphotyrosine antibody" refers to a molecule, usually an antibody, which binds selectively to phosphorylated tyrosine

residues in the kinase domain of a rPTK. The antibody can be polyclonal. but is desirably a monoclonal antibody. Anti-phosphotyrosine polyclonal antibodies can be made using the techniques disclosed in White and Backer, Methods in Enzymology 201:65-67 [1991] and monoclonal anti-phosphotyrosine antibodies can be obtained commercially from Upstate Biologicals, Inc. (UBI, Lake Placid, NY), for example.

The word "label" when used herein refers to a detectable compound or composition which is conjugated directly or indirectly with a molecule (such as the anti-phosphotyrosine antibody). The label may be detectable 10 by itself (e.g. radioisotope labels or fluorescent labels) or, in the case of an enzymatic label, may catalyze a chemical alteration of a substrate compound or composition which is detectable. The preferred label is an enzymatic one which catalyzes a color change of a non-radioactive color reagent.

By "washing" is meant exposing the solid phase to an aqueous solution (usually a buffer or cell culture media) in such a way that unbound material (e.g., non-adhering cells, non-adhering capture agent, unbound ligand, receptor, receptor construct, cell lysate, or anti-phosphotyrosine antibody) is removed therefrom. To reduce background noise, it is 20 convenient to include a detergent (e.g. Triton X) in the washing solution. Usually, the aqueous washing solution is decanted from the wells of the assay plate following washing. Conveniently, washing can be achieved using an automated washing device. Sometimes, several washing steps (e.g., between about 1 to 10 washing steps) may be required.

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By "block buffer" is meant an aqueous, pH buffered solution containing at least one blocking compound which is able to bind to exposed surfaces of the second solid phase which are not coated with capture agent. The blocking compound is normally a protein such as bovine serum albumin (BSA), gelatin, casein or milk powder and does not cross-react with any of the reagents in the assay (e.g., the anti-phosphotyrosine antibodies and detection reagents). The block buffer is generally provided at a pH between about 7 to 7.5 and suitable buffering agents include phosphate and TRIS.

By "lysis buffer" is meant an aqueous, pH buffered solution comprising a solubilizing detergent, one or more protease inhibitors and at least one phosphatase inhibitor (such as sodium orthovanadate). The term "solubilizing detergent" refers to a water miscible, non-ionic detergent which lyses cell membranes of eukaryotic cells but does not

denature or activate the receptor or receptor construct. Examples of suitable non-ionic detergents include Triton-X 100, Tween 20. CHAPS and Nonidet P-40 (NP40) available from Calbiochem, La Jolla, California, for example. Many other non-ionic detergents are available in the art. Examples of suitable protease inhibitors include phenylmethylsulfonyl fluoride (PMSF), leupeptin, pepstatin, aprotinin, 4-(2-aminoethyl)-benzenesulfonyl fluoride hydrochloride-bestatin, chymostatin and benzamidine. Preservatives (e.g., thimerosal) and one or more compounds which maintain the isotonicity of the solution (e.g., sodium chloride (NaCl) or sucrose) and a buffer (e.g., Tris or PBS) are usually also present. Generally, the pH of the lysis buffer is in the range about 7 to 7.5.

Usually, following addition of the lysis buffer to the first assay plate, the first assay plate is "gently agitated" and this expression refers to the act of physically shaking the first assay plate (normally using a circular motion) at a substantially low velocity. Gentle agitation does not involve mechanically disrupting the cells (e.g. by homogenizing or centrifuging the cells). Exemplary shaking velocities are in the order of 200 to 500 rpm, preferably 300 to 400 rpm in a Bellco orbital shaker, for example.

#### 20 II. Modes for Practicing the Invention

#### Kinase Receptor Activation - KIRA

The first stage of the assay involves phosphorylation of the kinase domain of a kinase receptor, wherein the receptor is present in the cell membrane of a eukaryotic cell. The receptor may be an endogenous receptor or nucleic acid encoding the receptor may be transformed into the cell. In one embodiment of the invention, nucleic acid encoding a receptor construct is transformed into the cell. Exemplary techniques for transforming the cell with either the receptor or the receptor construct nucleic acid follow.

#### A. Transformation of the cells

The instant invention provides a substantial improvement over soluble kinase receptor assays insofar as it is considered to more accurately reflect the activity of the receptor in situ. It has been discovered that it is possible to transform eukaryotic cells with a receptor construct (comprising the kinase receptor and either an amino- or carboxyl-terminal flag polypeptide) so that the receptor construct assembles itself appropriately in the cell membrane and still retains tyrosine kinase

activity which can be detected in the ELISA stage of the assay. This provides a generic assay for measuring tyrosine kinase activity of any tyrosin kinase of interest.

If a suitable capture agent as described herein is available for a 5 selected rPTK, cells can be transformed with the nucleic acid encoding the receptor alone, without the flag polypeptide. Alternatively, if cells are available which produce the receptor (e.g., MCF-7 cells which produce the HER2 receptor), it is not necessary to transform the cells for use in the assay.

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In order to transform the cells with the nucleic acid encoding the rPTK or receptor construct, nucleic acid encoding the rPTK and, optionally, the flag polypeptide, is isolated. This can be achieved by screening a cDNA or genomic library known to contain the DNA encoding the rPTK or flag polypeptide of interest with a selected labelled probe (e.g., an antibody 15 or oligonucleotide- probe) for the rPTK or flag polypeptide, using standard procedures as described in chapters 10-12 of Sambrook et al., Molecular Cloning: A Laboratory Manual (New York: Cold Spring Harbor Laboratory Press, 1989), for example. Alternatively, the nucleic acid encoding the flag polypeptide can be made synthetically using an oligo-synthesizing machine (Applied Biosystems, CA). An alternative means to isolate the nucleic acid encoding the rPTK or flag polypeptide is to use PCR methodology as described in section 14 of Sambrook et al., supra. Isolation of only the ECD of the rPTK of interest is required, since this nucleic acid can be fused to the nucleic acid encoding the TM and ICD of the Rse-flag polypeptide construct disclosed herein. See Figures 1A - 1C and SEQ ID NOS: 1 and 2. If necessary however, conventional primer extension procedures as described in section 7.79 of Sambrook et al., supra, can be used to detect precursors and processing intermediates of mRNA that may not have been reverse-transcribed into cDNA.

A preferred method of practicing this invention is to use carefully selected oligonucleotide sequences to screen cDNA libraries from various tissues, preferably mammalian cell lines having the rPTK of interest. The oligonucleotide sequences selected as probes should be of sufficient length and sufficiently unambiguous that false positives are minimized.

The oligonucleotide must be labeled such that it can be detected upon hybridization to DNA in the library being screened. The preferred method of labeling is to use 32P- labeled ATP with polynucleotide kinase, as is well known in the art, to radiolabel the oligonucleotide. However, other methods may be used to label the oligonucleotide, including, but not limited to, biotinylation or enzyme labeling.

In order to provide nucleic acid encoding a receptor construct, nucleic acid encoding the rPTK is fused at its 3' end to nucleic acid encoding the N-terminus of the flag polypeptide. Alternatively, the nucleic acid encoding the rPTK will be fused at its 5' end to nucleic acid encoding the carboxyl terminus of the flag polypeptide. Thus, the flag polypeptide is provided at either the carboxyl- or amino- terminus of the receptor construct. Examples of suitable flag polypeptides are provided above.

10 Selection of other suitable flag polypeptides is possible using the techniques described herein.

In order to generate fusions between the Rse.flag reagent and a rPTK of interest, the nucleic acid encoding the ECD of the rPTK of interest is fused at its 3' end to the nucleic acid encoding the amino terminus of the Rse.flag reagent.

The nucleic acid (e.g., cDNA or genomic DNA) encoding the rPTK or receptor construct is then inserted into a replicable vector for further cloning (amplification of the DNA) or for expression. Many vectors are available to the skilled practitioner but must be compatible with the cell which is to be used in the assay. The vector will have vector components the presence of which will depend on various factors. Such components include, for example, a signal sequence, an origin of replication, one or more marker genes, an enhancer element, a promoter, and a transcription termination sequence. Selection of these vector components shall be described below.

Incorporation of a signal sequence into the expression vector is required since the rPTK or receptor construct must be transported to the cell membrane where it is positioned such that the ECD faces the external milieu of the cell. Therefore, a signal sequence suitable for positioning the rPTK or receptor construct in such a manner is used. The signal sequence is generally a component of the vector, or it may be a part of the rPTK or receptor construct DNA that is inserted into the vector. If a heterologous signal sequence is used, it is from those that are recognized and processed (i.e., cleaved by a signal peptidase) by the host cell.

For yeast secretion the native signal sequence may be substituted by, e.g., the yeast invertase leader, alpha factor leader (including Saccharomyces and Kluyveromyces α-factor leaders, the latter described in U.S. Patent No. 5,010,182 issued 23 April 1991), or acid phosphatase

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leader, the *C. albicans* glucoamylase leader (EP 362,179 published 4 April 1990), or the signal described in WO 90/13646 published 15 November 1990. In mammalian cells expression of the DNA encoding the native signal sequence (e.g., the rPTK pre-sequence that normally directs secretion of rPTK from mammalian cells in vivo) is satisfactory, although other mammalian signal sequences may be suitable, such as signal sequences from other animal rPTKs, and signal sequences from secreted polypeptides of the same or related species, as well as viral secretory leaders, for example, the herpes simplex gD signal.

The DNA for such precursor region is ligated in reading frame to DNA encoding the rPTK or receptor construct.

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Both expression and cloning vectors contain a nucleic acid sequence that enables the vector to replicate in one or more selected host cells. Generally, in cloning vectors this sequence is one that enables the vector to replicate independently of the host chromosomal DNA, and includes origins of replication or autonomously replicating sequences. The 2µ plasmid origin is suitable for yeast, and various viral origins (SV40, polyoma, adenovirus, VSV or BPV) are useful for cloning vectors in mammalian cells. Generally, the origin of replication component is not needed for mammalian expression vectors (the SV40 origin may typically be used only because it contains the early promoter).

Most expression vectors are "shuttle" vectors, i.e., they are capable of replication in at least one class of organisms but can be transformed into another organism for expression. For example, a vector is cloned in E. coli and then the same vector is transformed into yeast or mammalian cells for expression even though it is not capable of replicating independently of the host cell chromosome.

DNA may also be amplified by insertion into the host genome. This is readily accomplished using Bacillus species as hosts, for example, by including in the vector a DNA sequence that is complementary to a sequence found in Bacillus genomic DNA. Transformation of Bacillus with this vector results in homologous recombination with the genome and insertion of rPTK or receptor construct DNA. However, the recovery of genomic DNA encoding the rPTK or receptor construct is more complex than that of an exogenously replicated vector because restriction enzyme digestion is required to excise the rPTK or receptor construct DNA.

Expression and cloning vectors usually contain a selection gene, also termed a selectable marker. This gene encodes a protein necessary for the

survival or growth of transformed host cells grown in a selective culture medium. Host cells not transformed with the vector containing the selection gene will not survive in the culture medium. Typical selection genes encode proteins that (a) confer resistance to antibiotics or other toxins, e.g., ampicillin, neomycin, methotrexate, or tetracycline, (b) complement auxotrophic deficiencies. or (c) supply critical nutrients not available from complex media.

One example of a selection scheme utilizes a drug to arrest growth of a host cell. Those cells that are successfully transformed with a 10 heterologous gene express the DNA encoding a protein conferring drug resistance and thus survive the selection regimen. Examples of such dominant selection use the drugs neomycin (Southern et al., J. Molec. Appl. Genet. 1:327 [1982]), mycophenolic acid (Mulligan et al., Science 209:1422 [1980]) or hygromycin (Sugden et al., Mol. Cell. Biol. 5:410-413 [1985]).

15 The three examples given above employ bacterial genes under eukaryotic control to convey resistance to the appropriate drug G418 or neomycin (geneticin), xgpt (mycophenolic acid), or hygromycin, respectively.

Another example of suitable selectable markers for mammalian cells are those that enable the identification of cells competent to take up the rPTK or receptor construct nucleic acid, such as DHFR or thymidine kinase. The mammalian cell transformants are placed under selection pressure that only the transformants are uniquely adapted to survive by virtue of having taken up the marker. Selection pressure is imposed by culturing the transformants under conditions in which the concentration of selection agent in the medium is successively changed, thereby leading to amplification of both the selection gene and the DNA that encodes the rPTK or receptor construct. Amplification is the process by which genes in greater demand for the production of a protein critical for growth are reiterated in tandem within the chromosomes of successive generations of 30 recombinant cells. Increased quantities of rPTK or receptor construct are synthesized from the amplified DNA. Other examples of amplifiable genes include metallothionein-I and -II, preferably primate metallothionein genes, adenosine deaminase, ornithine decarboxylase, etc.

For example, cells transformed with the DHFR selection gene are first identified by culturing all of the transformants in a culture medium that contains methotrexate (Mtx), a competitive antagonist of DHFR. An appropriate host cell when wild-type DHFR is employed is the Chinese hamster ovary (CHO) cell line deficient in DHFR activity, prepared and

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propagated as described by Urlaub and Chasin, Proc. Natl. Acad. Sci. USA 77:4216 (1980). The transformed cells are then exposed to increased levels of methotrexate. This leads to the synthesis of multiple copies of the DHPR gene, and, concomitantly, multiple copies of other DNA comprising the expression vectors, such as the DNA encoding the rPTK or receptor construct. This amplification technique can be used with any otherwise suitable host, e.g., ATCC No. CCL61 CHO-K1, notwithstanding the presence of endogenous DHFR if, for example, a mutant DHFR gene that is highly resistant to Mtx is employed (EP 117,060).

Alternatively, host cells (particularly wild-type hosts that contain endogenous DHFR) transformed or co-transformed with DNA sequences encoding the rPTK or receptor construct, wild-type DHFR protein, and another selectable marker such as aminoglycoside 3' phosphotransferase (APH) can be selected by cell growth in medium containing a selection agent for the selectable marker such as an aminoglycosidic antibiotic. e.g., kanamycin, neomycin, or G418. See U.S. Patent No. 4,965,199.

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A suitable selection gene for use in yeast is the trpl gene present in the yeast plasmid YRp7 (Stinchcomb et al., Nature 282:39 [1979]; Kingsman et al., Gene 7:141 [1979]; or Tschemper et al., Gene 10:157 [1980]). The trpl gene provides a selection marker for a mutant strain of yeast lacking the ability to grow in tryptophan, for example, ATCC No. 44076 or PEP4-1 (Jones, Genetics 85:12 [1977]). The presence of the trpl lesion in the yeast host cell genome then provides an effective environment for detecting transformation by growth in the absence of tryptophan. Similarly, Leu2-deficient yeast strains (ATCC 20,622 or 38,626) are complemented by known plasmids bearing the Leu2 gene.

In addition, vectors derived from the 1.6 µm circular plasmid pKD1 can be used for transformation of Kluyveromyces yeasts. Bianchi et al., Curr. Genet. 12:185 (1987). More recently, an expression system for large-scale production of recombinant calf chymosin was reported for K. lactis. Van den Berg, Bio/Technology 8:135 (1990). Stable multi-copy expression vectors for secretion of mature recombinant human serum albumin by industrial strains of Kluyveromyces have also been disclosed. Pleer et al., Bio/Technology 9:968-975 (1991).

Expression and cloning vectors usually contain a promoter that is recognized by the host organism and is operably linked to the rPTK or receptor construct nucleic acid. Promoters are untranslated sequences located upstream (5') to the start codon of a structural gene (generally

within about 100 to 1000 bp) that control the transcription and translation of particular nucleic acid sequence, such as the rPTK nucleic acid sequence, to which they are operably linked. Such promoters typically fall into two classes, inducible and constitutive. Inducible promoters are 5 promoters that initiate increased levels of transcription from DNA under their control in response to some change in culture conditions, e.g., the presence or absence of a nutrient or a change in temperature. At this time a large number of promoters recognized by a variety of potential host cells are well known. These promoters are operably linked to rPTK or receptor 10 construct-encoding DNA by removing the promoter from the source DNA by restriction enzyme digestion and inserting the isolated promoter sequence into the vector. Both the native rPTK promoter sequence and many heterologous promoters may be used to direct amplification and/or expression of the rPTK or receptor construct DNA. The promoter will be one which results in the accumulation of suitable numbers of receptor or receptor construct in the cell membrane of the transformed cell (i.e. so that autophosphorylation of the receptor is detectable in the ELISA but constitutive phosphorylation does not occur). Selection of a suitable promoter to achieve this is possible following the guidelines herein for 20 selecting cells for use in the KIRA ELISA.

Virtually all eukaryotic genes have an AT-rich region located approximately 25 to 30 bases upstream from the site where transcription is initiated. Another sequence found 70 to 80 bases upstream from the start of transcription of many genes is a CXCAAT region where X may be any nucleotide. At the 3' end of most eukaryotic genes is an AATAAA sequence that may be the signal for addition of the poly A tail to the 3' end of the coding sequence. All of these sequences are suitably inserted into eukaryotic expression vectors.

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Examples of suitable promoting sequences for use with yeast hosts include the promoters for 3-phosphoglycerate kinase (Hitzeman et al., J. Biol. Chem. 255:2073 [1980]) or other glycolytic enzymes (Hess et al., J. Adv. Enzyme Reg., 7:149 [1968]; and Holland, Biochemistry 17:4900 [1978]), such as enolase, glyceraldehyde-3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglucose isomerase, and glucokinase.

Other yeast promoters, which are inducible promoters having the additional advantage of transcription controlled by growth conditions, are

the promoter regions for alcohol dehydrogenase 2, isocytochrome C. acid phosphatase, degradative enzymes associated with nitrogen metabolism, metallothionein, glyceraldehyde-3-phosphate dehydrogenase, and enzymes responsible for maltose and galactose utilization. Suitable vectors and promoters for use in yeast expression are further described in Hitzeman et al., EP 73,657A. Yeast enhancers also are advantageously used with yeast promoters.

rPTK or receptor construct transcription from vectors in mammalian host cells is controlled, for example, by promoters obtained from the genomes of viruses such as polyoma virus, fowlpox virus (UK 2,211,504 published 5 July 1989), adenovirus (such as Adenovirus 2), bovine papilloma virus, avian sarcoma virus, cytomegalovirus, a retrovirus, hepatitis-B virus and most preferably Simian Virus 40 (SV40), from heterologous mammalian promoters, e.g., the actin promoter or an immunoglobulin promoter, from heat-shock promoters, and from the promoter normally associated with the rPTK or receptor construct sequence, provided such promoters are compatible with the host cell systems.

The early and late promoters of the SV40 virus are conveniently obtained as an SV40 restriction fragment that also contains the SV40 viral origin of replication. Fiers et al., Nature 273:113 (1978); Mulligan and Berg, Science 209:1422-1427 (1980); Pavlakis et al., Proc. Natl. Acad. Sci. The immediate early promoter of the human <u>USA</u> 78:7398-7402 (1981). cytomegalovirus is conveniently obtained as a HindIII E restriction fragment. Greenaway et al., Gene 18:355-360 (1982). A system for expressing DNA in mammalian hosts using the bovine papilloma virus as a vector is disclosed in U.S. 4,419,446. A modification of this system is described in U.S. 4,601,978. See also Gray et al., Nature 295:503-508 (1982) on expressing cDNA encoding immune interferon in monkey cells; Reyes et al., Nature 297:598-601 (1982) on expression of human  $\beta$ -interferon cDNA in mouse cells under the control of a thymidine kinase promoter from herpes simplex virus; Canaani and Berg, Proc. Natl. Acad. Sci. USA 79:5166-5170 (1982) on expression of the human interferon \$1 gene in cultured mouse and rabbit cells; and Gorman et al., Proc. Natl. Acad. Sci. USA 79:6777-6781 (1982) on expression of bacterial CAT sequences in CV-1 monkey kidney cells, chicken embryo fibroblasts, Chinese hamster ovary cells, HeLa cells, and mouse NIH-3T3 cells using the Rous sarcoma virus long terminal repeat as a promoter.

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Transcription of DNA encoding the rPTK or receptor construct by higher eukaryotes may be increased, if increased numbers of the rPTK or receptor construct per cell are required to facilitate detection in the ELISA stage of the assay. This may be achieved by inserting an enhancer sequence into the vector. Enhancers are cis-acting elements of DNA, usually about from 10 to 300 bp, that act on a promoter to increase its Enhancers are relatively orientation and position transcription. independent, having been found 5' (Laimins et al., Proc. Natl. Acad. Sci. USA 78:993 [1981]) and 3' (Lusky et al., Mol. Cell Bio. 3:1108 [1983]) to the transcription unit, within an intron (Banerji et al., Cell 33:729 [1983]), as well as within the coding sequence itself (Osborne et al., Mol. Cell Bio. 4:1293 [1984]). Many enhancer sequences are now known from mammalian genes (globin, elastase, albumin, a-fetoprotein, and insulin). Typically, however, one will use an enhancer from a eukaryotic cell virus. Examples include the SV40 enhancer on the late side of the replication origin (bp 100-270), the cytomegalovirus early promoter enhancer, the polyoma enhancer on the late side of the replication origin, and adenovirus enhancers. See also Yaniv, Nature 297:17-18 (1982) on enhancing elements for activation of eukaryotic promoters. The enhancer may be spliced into 20 the vector at a position 5' or 3' to the rPTK or receptor constructencoding sequence, but is preferably located at a site 5' from the promoter.

Expression vectors used in eukaryotic host cells will also contain sequences necessary for the termination of transcription and for stabilizing the mRNA. Such sequences are commonly available from the 5' and, occasionally 3', untranslated regions of eukaryotic or viral DNAs or cDNAs. These regions contain nucleotide segments transcribed as polyadenylated fragments in the untranslated portion of the mRNA encoding the rPTK or receptor construct.

Construction of suitable vectors containing one or more of the above listed components employs standard ligation techniques. Isolated plasmids or DNA fragments are cleaved, tailored, and religated in the form desired to generate the plasmids required.

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For analysis to confirm correct sequences in plasmids constructed, the ligation mixtures are used to transform E. coli K12 strain 294 (ATCC 31,446) and successful transformants selected by ampicillin or tetracycline resistance where appropriate. Plasmids from the transformants are prepared, analyzed by restriction endonuclease digestion, and/or sequenced

by the method of Messing et al., Nucleic Acids Res. 9:309 (1981) or by the method of Maxam et al., Methods in Enzymology 65:499 (1980).

Other methods, vectors, and h st cells suitable for adaptation to the synthesis of the rPTK or receptor construct in recombinant vertebrate cell culture are described in Gething et al., Nature 293:620-625 (1981); Mantei et al., Nature 281:40-46 (1979); Levinson et al.; EP 117.060; and EP A particularly useful plasmid for mammalian cell culture 117,058. expression of rPTK or receptor construct DNA is pRK5 (EP 307,247) or pSVI6B (PCT pub. no. WO 91/08291 published 13 June 1991).

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Examples of suitable eukaryotic cell lines for transformation include Saccharomyces cerevisiae, Schizosaccharomyces pombe (Beach and Nurse, Nature 290:140 [1981]; EP 139,383 published 2 May 1985); Kluyveromyces hosts (U.S. 4,943,529; Fleer et al., Bio/Technology 9:968-975 [1991]) and Aspergillus hosts such as A. nidulans (Ballance et al., Biochem. Biophys. 15 Res. Commun. 112:284-289 [1983]; Tilburn et al., Gene 26:205-221 [1983]; Yelton et al., Proc. Natl. Acad. Sci. USA 81:1470-1474 [1984]) and A. niger (Kelly and Hynes, EMBO J. 4:475-479 [1985]), among lower eukaryotic host microorganisms.

Examples of useful animal host cell lines for transformation include monkey kidney CV1 line transformed by SV40 (COS-7, ATCC CRL 1651); human embryonic kidney line (293 or 293 cells subcloned for growth in suspension culture, Graham et al., J. Gen Virol. 36:59 [1977]); baby hamster kidney cells (BHK, ATCC CCL 10); Chinese hamster ovary cells/-DHFR (CHO, Urlaub and Chasin, Proc. Natl. Acad. Sci. USA 77:4216 [1980]); mouse sertoli cells (TM4, Mather, Biol. Reprod. 23:243-251 [1980]); monkey kidney cells (CV1 ATCC CCL 70); African green monkey kidney cells (VERO-76, ATCC CRL-1587); human cervical carcinoma cells (HELA, ATCC CCL 2); canine kidney cells (MDCK, ATCC CCL 34); buffalo rat liver cells (BRL 3A, ATCC CRL 1442); human lung cells (W138, ATCC CCL 75); human liver cells (Hep G2, HB 8065); mouse 30 mammary tumor (MMT 060562, ATCC CCL51); TRI cells (Mather et al., Annals N.Y. Acad. Sci. 383:44-68 [1982]); MRC 5 cells; FS4 cells; and a human hepatoma line (Hep G2).

Host cells are transformed with the above-described expression or cloning vectors of this invention and cultured in conventional nutrient media modified as appropriate for inducing promoters, selecting transformants, or amplifying the genes encoding the desired sequences. Transformation means introducing DNA into an organism so that the DNA is replicable, either as an extrachromosomal element or as a chromosomal

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integrant. Depending on the host cell used, transformation is done using standard techniques appropriate to such cells. Successful transformation is generally recognized when any indication of the operation of this vector occurs within the host cell.

For mammalian cells, the calcium phosphate precipitation method of Graham and Van der Eb, Virology 52:456-457 (1978) is preferred. General aspects of mammalian cell host system transformations have been described by Axel in U.S. 4,399,216 issued 16 August 1983. Transformations into yeast are typically carried out according to the method of Van Solingen et 10 al., J. Bact. 130:946 (1977) and Hsiao et al., Proc. Natl. Acad. Sci. (USA) 76:3829 (1979). However, other methods for introducing DNA into cells, such as by nuclear microinjection, electroporation, bacterial protoplast fusion with intact cells, or polycations, e.g., polybrene, polyornithine, etc., may also be used. For various techniques for transforming mammalian 15 cells, see Keown et al., Methods in Enzymology (1989), Keown et al., Methods in Enzymology 185:527-537 (1990), and Mansour et al., Nature 336:348-352 (1988).

The mammalian host cells used to produce the rPTK or receptor construct may be cultured in a variety of media. Commercially available media such as Ham's F10 (Sigma), Minimal Essential Medium ([MEM], Sigma), RPMI-1640 (Sigma), and Dulbecco's Modified Eagle's Medium ([DMEM], Sigma) are suitable for culturing the host cells. In addition, any of the media described in Ham and Wallace, Meth. Enz. 58:44 (1979), Barnes and Sato, Anal. Biochem. 102:255 (1980). U.S. 4,767,704; 4,657,866; 4,927,762; or 4,560,655; WO 90/03430; WO 87/00195; U.S. Patent Re. 30,985; or U.S. Patent No. 5,122,469, the disclosures of each of which are incorporated herein by reference, may be used as culture media for the host cells. Any of these media may be supplemented as necessary with hormones and/or other growth factors (such as insulin, transferrin, or epidermal growth factor), salts 30 (such as sodium chloride, calcium, magnesium, and phosphate), buffers (such as HEPES), nucleosides (such as adenosine and thymidine), antibiotics (such as Gentamycin<sup>M</sup> drug), trace elements (defined as inorganic compounds usually present at final concentrations in the micromolar range), and glucose or an equivalent energy source. Any other necessary supplements 35 may also be included at appropriate concentrations that would be known to those skilled in the art. The culture conditions, such as temperature, pH, and the like, are those previously used with the host cell selected for expression, and will be apparent to the ordinarily skilled artisan.

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In general, principles, protocols, and practical techniques for maximizing the productivity of mammalian cell cultures can be found in Mammalian Cell Biotechnology: a Practical Approach, M. Butler, ed., IRL Press, 1991.

Gene amplification and/or expression may be measured in a sample directly, for example, by conventional Southern blotting, Northern blotting to quantitate the transcription of mRNA (Thomas, Proc. Natl. Acad. Sci. USA 77:5201-5205 [1980]), dot blotting (DNA analysis), or in situ hybridization, using an appropriately labeled probe, based on the sequences Various labels may be employed, most commonly 10 provided herein. radioisotopes, particularly 32P. However, other techniques may also be employed, such as using biotin-modified nucleotides for introduction into a polynucleotide. The biotin then serves as the site for binding to avidin or antibodies, which may be labeled with a wide variety of labels, such as 15 radionuclides, fluorophores, enzymes, or the like. Alternatively, antibodies may be employed that can recognize specific duplexes, including DNA duplexes, RNA duplexes, and DNA-RNA hybrid duplexes or DNA-protein duplexes. The antibodies in turn may be labeled and the assay may be carried out where the duplex is bound to a surface, so that upon the 20 formation of duplex on the surface, the presence of antibody bound to the duplex can be detected.

Gene expression, alternatively, may be measured by immunological methods, such as immunohistochemical staining to quantitate directly the expression of gene product.

#### B. Selecting cells for use in the assay

As mentioned above, the cells to be subjected to the assay can be (a) cells having an endogenous receptor, (b) cells which have been transformed with a rPTK, or (c) cells transformed with a receptor construct. suitability of the cells for use in the assay is investigated.

Cells having the endogenous rPTK can be subjected to a test-run KIRA ELISA using a known ligand to the PTK (e.g. an agonist antibody) and a control (e.g. the diluent for the agonist antibody). A range of ligand concentrations such as those used herein (see Examples 1, 2 and 3) will be used to determine whether sufficient numbers of the receptor are present in the cells being tested. In order to discover whether a cell line is unsuitable because the receptor is constitutively phosphorylated, the cell line can be subjected to the KIRA ELISA disclosed herein, wherein it is exposed to both positive and negative controls (e.g. a known agonist ligand in cell culture media as described herein as a positive control and the cell culture media without the agonist ligand as the negative control). If phosphorylation of the receptor is detected for both positive and negative controls, this may be indicative that constitutive phosphorylation of the receptor is occurring. However, it is possible that a constituent of the serum in the cell culture media is activating the receptor. Thus, the cells can be "starved" in serum-free media for about 2-12 hours (depending on cell survival) and then the assay is repeated using the positive and negative controls. If activation is detected for both controls, the cell line may be considered unsuitable and another cell line can be tested.

If the cell line is transformed with the receptor (without the flag polypeptide) a strategy similar to that depicted in Figure 4 can be used to discover whether or not the cell line is suitable for use in the assay. 15 As a first step, successful transformation and expression of the nucleic acid encoding the rPTK is determined (see Figure 4, step b). In order to identify whether the ECD of the rPTK is present on the surface of the cells, flow cytometric analysis can be performed using an antibody to the ECD of the receptor. The antibody can be made using the techniques for generating antibodies discussed herein. Flow cytometric analysis can be carried out using the techniques described in Current Protocols in Immunology, Ed. Coligen et al., Wiley publishers, Vols. 1 and 2, for example. Briefly, flow cytometric analysis involves incubating intact cells (having the receptor) with antibodies to the ECD thereof, followed by washing. The antibody-bound cells are then incubated with species specific anti-antibody antibodies conjugated to a fluorochrome. Following washing, the labeled cells are analyzed by fluorescence-activated flow cytometry to detect whether the ECD is present on the surface of the cells.

In the following step, i.e. Figure 4, step (c), the ability of the cell-bound receptor to be activated is tested. In order to determine this, the transformed cells are exposed to a known agonist to the receptor (e.g. the endogenous ligand or an agonist antibody for the receptor). Following exposure, the cells are lysed in a suitable buffer (e.g. sodium dodecylbenzenesulfonate in phosphate buffered saline; SDS in PBS) and subjected to Western blotting with anti-phosphotyrosine antibodies as described in Wang, Molecular and Cellular Biology 5(12):3640-3643 (1985); Glenney et al., Journal of Immunological Methods 109:277-285 (1988); Kamps, Methods in Enzymology 201:101-110 (1991); Kozma et al., Methods in

Enzymology 201:28-43 (1991); Holmes et al., Science 256:1205-10 (1992); or Corfas et al., PNAS. USA 90:1624-1628 (1993). for example.

Assuming the Western blotting step indicates that the rPTK can be activated, a KIRA ELISA test run can be performed, see Figure 4 step (d), to further establish whether or not the transformed cell line can be used in the assay.

In the preferred embodiment of the invention, the KIRA ELISA is a "generic" assay insofar as any rPTK of interest can be studied regardless of the availability of receptor-specific reagents (i.e., capture agent). This embodiment employs a receptor construct having a flag polypeptide at either the amino or carboxyl terminus of the receptor.

If the flag polypeptide is provided at the NH<sub>2</sub>-terminus (see, e.g., the gD.trk A, B and C receptor constructs disclosed in Example 3), the procedure for selecting a transformed cell line for use in the assay summarized in Figure 4 can be performed. In this embodiment, the cells are transformed with the flag polypeptide-receptor construct as described earlier herein. See step (a). In step (b), successful transformation of the receptor and flag polypeptide (i.e. the receptor construct) is confirmed. In order to study this, two-dimensional flow cytometric analysis can be performed using antibodies to both the flag polypeptide and the ECD of the receptor. Techniques for two-dimensional flow cytometric analysis are disclosed in Current Protocols in Immunology, supra. Assuming successful transformation of the receptor construct is demonstrated, steps (c) and (d) of Figure 4 are then performed. See the discussion above, for an explanation of steps (c) to (d) of Figure 4.

A technique for identification of cells which have been successfully transformed with the receptor construct having a C-terminal flag polypeptide and which cells are also suitable for use in the assay is illustrated in Figure 5. Following cell transformation [step (a)], successful transformation of the receptor is determined by flow cytometric analysis using an antibody directed against the ECD of the receptor of interest, for example. Flow cytometric analysis can be performed substantially as described above. This forms step (b) of the procedure outlined in Figure 5.

Following step (b), successful transformation of the entire receptor construct (including the COOH-terminal flag polypeptide) is analyzed in step (c). This can be achieved by lysing the cells (using techniques for lysing cells disclosed herein) and immunoprecipitating the membrane extract

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with an antibody against the receptor of interest. This immunoprecipitated membrane extract is then subjected to Western blot analysis with antibodies specific for the flag polypeptide. Alternatively, rPTK-specific ELISA analysis of anti-flag polypeptide captured membrane lysate can be carried 5 out. Briefly, this involves coating ELISA wells with appropriate flag specific capture agent. The wells are blocked, washed, and the lysate is then incubated in the wells. Unbound receptor construct is removed by washing. The wells are then reacted with receptor-specific antibody or antibodies, either directly or indirectly conjugated to HRPO. The wells 10 are washed and the HRPO is then exposed to the chromogenic substrate (e.g., TMB).

Steps (d) and (e), i.e., detecting receptor activation and KIRA ELISA test run, are essentially the same as those steps described above.

Once useful cells are identified, they are subjected to the KIRA 15 stage of the instantly claimed assay.

C. Coating the first solid phase with the cells

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The first solid phase (e.g. a well of a first assay plate) is coated with cells having the endogenous receptor or cells which have been transformed pursuant to the preceding sections.

Preferably, an adherent cell line is chosen, so that the cells naturally adhere to the first solid phase. However, use of an adherent cell line is not essential. For example, non-adherent cells (e.g. red blood cells) can be added to round bottomed wells of an assay plate such as that sold by Becton Dickinson Labware, Lincoln Park, New Jersey, for example. The assay plate is then placed in a plate carrier and centrifuged so as to create a pellet of cells adhering to the base of the wells. culture supernatants are removed using a pipette. Thus, use of an adherent cell is clearly advantageous over non-adherent cells since it reduces variability in the assay (i.e, the cells in the pellet of the round bottom 30 wells may be taken up with the supernatant when the alternative method is used).

The cells to be added to the wells of the first assay plate may be maintained in tissue culture flasks and utilized when cells densities of about 70-90% confluency are achieved. Then, generally between about 1 x 35 104 to 3 x 105 (and preferably 5 x 104 to 1 x 105) cells are seeded per flat-bottom well, using a pipette, for example. It has been found that, contrary to expectations, addition of cell concentrations mentioned above is sufficient to enable activation of the rPTK to be measured in the ELISA 5

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stage of the assay, without the need to concentrate or clarify the cells or cell lysate prior thereto. Often, the cells are diluted in culture medium pri r to seeding them in the wells of the microtiter plate to achieve the desired cell densities.

Usually, the cells are cultured in the microtiter plates for a sufficient period of time to optimize adherence to the wells thereof. but not too long such that the cells begin to deteriorate. Thus, incubation for about 8 to 16 hours at a temperature which is the physiological optimum for the cells (usually about 37°C) is preferred. Suitable media for 10 culturing the cells are described in Section 1A above. Culturing in 5% CO2 is recommended.

Following incubation overnight, the well supernatants are decanted and excess supernatant may be further removed by lightly tamping the microtiter plates with an absorbent substrate, e.g., a paper towel, but a 15 sponge works equally well. Thus, a substantially homogeneous layer of adhering cells remains on the internal surfaces of the individual wells of the microtiter plate. These adhering cells are then exposed to the analyte.

# D. Preparation and addition of the analyte

As mentioned above, the analyte may comprise an agonist ligand (or suspected agonist) or an antagonist (or suspected antagonist) for the rPTK of interest. The ligand may be an endogenous polypeptide, or a synthetic molecule, such as an inorganic or organic molecule. Usually, the ligand is a polypeptide. This assay is useful for screening molecules which activate (or antagonize activation) of the tyrosine kinase receptor of interest. Thus, the assay can be used for developing therapeutically effective molecules.

Where the ligand is an agonist, the molecule can comprise the native growth factor e.g., heregulin (HRG), insulin, insulin-like growth factors 30 I and II (IGF-I and IGF-II), epidermal growth factor (EGF), interleukins (e.g., IL-8), macrophage colony-stimulating factor (M-CSF), erythropoietin (EPO), platelet-derived growth factor (PDGF), transforming growth factors alpha and beta (TGF- $\alpha$  and TGF- $\beta$ ), hepatocyte growth factor (HGF), fibroblast growth factor (FGF) and nerve growth factor (NGF). Many of these growth factors are available commercially. Alternatively, the growth factor can be made by peptide synthesis or recombinant techniques which are described herein. Synthetic small molecule agonists can similarly be generated by those skilled in the art using conventional chemical synthesis techniques.

Where the ligand is present in a biological fluid, the analyte can be prepared using techniques which are well known in the art. Body fluid such as blood or ammiotic fluid may be used directly, however concentration may be required. If the analyte to be tested comprises a particular tissue, the cells thereof can be grown in cell culture and the supernatant can be tested for secreted ligand.

Often, the ligand is diluted in an aqueous diluent (such as cell culture media) so that a standard curve can be generated. However, the ligand may be present in a cell or a cell component (e.g., the cell membrane). In particular, it has been found that the assay can be used to detect the presence of a ligand in the cell membrane of a selected cell line. This is clearly useful for discovering a novel endogenous ligand for a known rPTK.

The ligand composition is added to each well which contains the adhering cells using a pipette, for example. At least one control well (e.g. to which the aqueous diluent for the ligand is added) is included in the assay.

The adhering cells are usually stimulated for a sufficient period of time to optimize the signal, but not too long such that the signal decreases as a consequence of dephosphorylation of the rPTK by endogenous phosphatases. A suitable stimulation period is between about 10 to 60 minutes, preferably about 30 minutes at a physiologically optimal temperature for the cells (usually about 37°C).

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Following activation, well supernatants are decanted and the plates can then be lightly tamped with an absorbent substrate to remove excess supernatant.

The assay can be used to detect antagonist ligands for the rPTK of interest. Antagonists generally fall into two categories (a) ones which bind to the rPTK and thereby block binding and/or activation of the rPTK by an agonist thereto (the antagonist may bind to the ECD, but this is not necessarily the case) and (b) those which bind to the agonist and thus prevent activation of the rPTK by the agonist.

In order to detect antagonist molecules from category (a) above, the cells are exposed to the suspected antagonist ligand substantially as mentioned above. Following exposure to the antagonist, the well supernatants are decanted and the plates are lightly tamped. Then, a known

agonist (e.g., the endogenous growth factor) is added to the washed cells essentially as discussed in the preceding paragraphs, following which, the well supernatants are decanted and plats are lightly tamped. Alternatively, a composition comprising both the antagonist and agonist can be added to the adhering cells substantially as discussed above. Ability of the suspected antagonist to block binding and/or activation of the rPTK can subsequently be measured by ELISA as discussed below.

To detect antagonist molecules from category (b) above, a known agonist is pre-incubated with the suspected antagonist prior to the KNRA stage of the assay. This incubation is carried out for a sufficient period of time to enable a complex of the antagonist-agonist to form; from 30 min. to 12 hours, for example. This complex is then subjected to the assay with the non-complexed agonist and antagonist used as controls.

Following exposure to the agonist (and optionally the antagonist)

5 ligand, the cells are lysed, as discussed below.

### E. Solubilizing the cells

In this step of the assay, the cells are lysed so as to solubilize the rPTK such that it remains activated (i.e., the tyrosine residues remain phosphorylated) for the ELISA stage of the assay. Thus, the cells are lysed using a lysis buffer as described above which serves to solubilize the rPTK or receptor construct, yet does not dephosphorylate or denature the rPTK.

Where microtiter plates are used as mentioned above, about 75 to 200µl of lysis buffer is added to each well. The plates can then be agitated gently using a plate shaker (e.g., such as that sold by Bellco Instruments, Vineland, NJ) for about 1 to 2 hours. Shaking can be carried out at room temperature.

# Enzyme-Linked Immunosorbent Assay - ELISA

The second stage of the assay involves a sandwich ELISA performed in the second assay plate. In order to carry out the ELISA, a capture agent is prepared.

# A. Preparation of the capture agent

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As mentioned above, the capture agent often comprises a polyclonal antibody (usually an affinity purified polyclonal antibody) or monoclonal antibody. Other capture agents are envisaged and are discussed in the definitions section above. The capture agent either binds specifically to the kinase receptor, or to the flag polypeptide (i.e. the antigen).

Polyclonal antibodies to the antigen (either the receptor or the flag polypeptide) generally are raised in animals by multiple subcutaneous (sc)

or intraperitoneal (ip) injections of the antigen or an antigenic fragment thereof (often the ECD of the rPTK) and an adjuvant. It may be useful to conjugate the antigen or a fragment containing the target amino acid sequence to a protein that is immunogenic in the species to be immunized (e.g., keyhole limpet hemocyanin), using a bifunctional or derivatizing agent.

The route and schedule for administration of immunogen to the host animal or cultured antibody-producing cells therefrom are generally in keeping with established and conventional techniques for antibody stimulation and production. While mice are frequently employed as the test model, it is contemplated that any mammalian subject including human subjects or antibody-producing cells obtained therefrom can be manipulated according to the processes of this invention to serve as the basis for production of mammalian, including human, hybrid cell lines.

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Animals are typically immunized against the immunogenic conjugates or derivatives by combining 1 mg or 1  $\mu$ g of conjugate (for rabbits or mice, respectively) with 3 volumes of Freund's complete adjuvant and injecting the solution intradermally at multiple sites. One month later the animals are boosted with 1/5 to 1/10 the original amount of conjugate in Freund's complete adjuvant (or other suitable adjuvant) by subcutaneous injection at multiple sites. 7 to 14 days later animals are bled and the serum is assayed for anti-antigen titer. Animals are boosted until the titer plateaus. Preferably, the animal is boosted with the conjugate of the same antigen, but conjugated to a different protein and/or through a different cross-linking agent. Conjugates also can be made in recombinant cell culture as protein fusions. Also, aggregating agents such as alum are used to enhance the immune response.

After immunization, monoclonal antibodies can be prepared by recovering immune cells (typically spleen cells or lymphocytes from lymph node tissue) from immunized animals and immortalizing the cells in conventional fashion, e.g., by fusion with myeloma cells or by Epstein-Barr (EB)-virus transformation and screening for clones producing the desired antibody. The hybridoma technique described originally by Kohler and Milstein, Eur. J. Immunol. 6:511 (1976), and also described by Hammerling et al., In: Monoclonal Antibodies and T-Cell Hybridomas, Elsevier, N.Y., pp. 563-681 (1981) has been widely applied to produce hybrid cell lines that secrete high levels of monoclonal antibodies against many specific antigens.

It is possible to fuse cells of one species with another. However, it is preferable that the source of the immunized antibody producing cells and the myeloma be from the same species.

The hybrid cell lines can be maintained in culture in cell culture media. The cell lines of this invention can be selected and/or maintained composition comprising the continuous cell line in hypoxanthine-aminopterin-thymidine (HAT) medium. In fact, once the hybridoma cell line is established, it can be maintained on a variety of nutritionally adequate media. Moreover, the hybrid cell lines can be 10 stored and preserved in any number of conventional ways, including freezing and storage under liquid nitrogen. Frozen cell lines can be revived and cultured indefinitely with resumed synthesis and secretion of monoclonal antibody.

The secreted antibody is recovered from tissue culture supernatant

by conventional methods such as precipitation, ion exchange chromatography,
affinity chromatography, or the like. The antibodies described herein are
also recovered from hybridoma cell cultures by conventional methods for
purification of IgG or IgM, as the case may be, that heretofore have been
used to purify these immunoglobulins from pooled plasma, e.g., ethanol or

polyethylene glycol precipitation procedures. The purified antibodies are
then sterile filtered. Where the antibody is a polyclonal antibody, it is
generally affinity purified using an affinity column generated from the
antigen of interest so as to provide a substantially specific capture
antibody. Affinity chromatography is usually preceded by other purification
techniques, such as liquid chromatography.

In a further embodiment, antibodies or antibody fragments can be isolated from antibody phage libraries generated via the techniques described in McCafferty et al., Nature, 348:552-554 (1990), using the flag polypeptide, rPTK, or a fragment thereof, to select for a suitable antibody or antibody fragment. Clackson et al., Nature, 352:624-628 (1991) and Marks et al., J. Mol. Biol., 222:581-597 (1991) describe the isolation of murine and human antibodies, respectively, using phage libraries. Subsequent publications describe the production of high affinity (nM range) human antibodies by chain shuffling (Mark et al., Bio/Technol. 10:779-783 [1992]), as well as combinatorial infection and in vivo recombination as a strategy for constructing very large phage libraries (Waterhouse et al., Nuc. Acids Res., 21:2265-2266 [1993]). Thus, these techniques are viable alternatives to traditional monoclonal antibody hybridoma techniques for

isolation of "monoclonal" antibodies which are encompassed by the present invention.

DNA encoding the monoclonal antibodies of the invention is readily isolated and sequenced using conventional procedures (e.g., by using 5 oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of murine antibodies). The hybridoma cells of the invention serve as a preferred source of such DNA. Once isolated, the DNA may be placed into expression vectors, which are then transfected into host cells such as simian COS cells, Chinese hamster ovary 10 (CHO) cells, or myeloma cells that do not otherwise produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. The DNA also may be modified, for example, by substituting the coding sequence for human heavy and light chain constant domains in place of the homologous murine sequences, Morrison et al., Proc. Nat. Acad. Sci. 81, 6851 (1984), or by covalently joining to the immunoglobulin coding sequence all or part of the coding sequence for a non-immunoglobulin polypeptide. In that manner, "chimeric" or "hybrid" antibodies are prepared that have the binding specificity of an anti-rPTK or anti-flag polypeptide monoclonal antibody herein. Thus, the antibody may be made by recombinant DNA methods (Cabilly et al., U.S. Pat. No. 4,816,567).

Binding of the capture agent is not affected by the presence or absence of a ligand bound to the receptor and the capture agent does not sterically block access to the phosphorylated tyrosine(s) by the antiphosphotyrosine antibody. Furthermore, the capture agent does not, of course, activate the receptor of interest. In order to screen for an antibody having these characteristics, the procedure outlined in Figure 3 can be carried out.

First, once the capture agent (e.g. an antibody or streptavidin) has

been chosen, binding to either the receptor or the flag polypeptide (where
a receptor construct is to be used in the assay) is confirmed. This can
be determined by flow cytometric analysis, immuno-precipitation or antigencoat ELISA, for example. Flow cytometric analysis has been described
above. Immunoprecipitation usually involves lysing the cells (having the
receptor or receptor construct) in non-ionic detergent (e.g. 0.5% Triton
X-100) in a suitable buffer (e.g. PBS) and the cell lysates thus obtained
are then incubated with the potential anti-receptor or anti-flag
polypeptide capture agent. The immune complexes are precipitated with

either (a) anti-capture agent antibodies in the presence of polyethylene glycol (PEG) which enhances precipitation of the immune complex or with (b) insoluble (e.g. agarose bound) protein A or protein G. The immunoprecipitated material is then analyzed by polyacrylamide gel electrophoresis (PAGE). For antigen-coat ELISA, ELISA wells are coated overnight with either the purified receptor, purified flag polypeptide or purified receptor construct. The coated wells are then exposed to the potential capture agent and screened with HRPO-conjugated species specific anti-capture agent antibody.

The ability of the capture agent to bind to the receptor or flag polypeptide in the presence of a ligand to the receptor is also confirmed. This can be analyzed by incubating the receptor or receptor construct with a known ligand for the receptor (e.g. the endogenous growth factor or an agonist antibody thereto). Flow cytometric analysis, immunoprecipitation 15 or antigen-coat ELISA can then be performed substantially as described above to investigate binding of the capture agent.

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Assuming the capture agent is suitable as determined by the preceding two steps, it is then shown that the capture agent does not induce receptor activation (i.e. autophosphorylation) either before or after cell lysis. Thus, the cell-bound receptor or receptor construct is exposed to either the potential capture agent or a negative control (e.g. a control antibody which does not activate the receptor). Following cell lysis, the receptor or receptor construct can be subjected to Western blot analysis using labeled anti-phosphotyrosine antibodies. See, e.g., Glenney et al., Journal of Immunological Methods 109:277-285 (1988); Kamps, Methods in Enzymology 201:101-110 (1991); Kozma et al., Methods in Enzymology 201:28-43 (1991); or Holmes et al., Science 256:1205-10 (1992). To establish whether the capture agent induces receptor activation following cell lysis, a trial run of the KIRA ELISA (with both the capture agent and a negative 30 control as discussed above) can be performed.

Finally, the ability of an anti-phosphotyrosine antibody (e.g. biotinylated anti-phosphotyrosine antibody) to bind the activated receptor in the presence of the potential capture agent is confirmed by a trial run in the KIRA ELISA disclosed herein.

Assuming the capture agent meets all the criteria specified above, it has good potential for use in the KIRA ELISA.

Once a suitable capture agent has been prepared, the second solid phase is coated therewith. Between about 0.1 to 10  $\mu$ g/ml of capture agent

can be added to each well of the second assay plate using a pipette, for example. The capture agent is often provided in a buffer at a high pH (e.g., between about 7.5 to 9.6) so that it has an increased overall charge and therefore displays enhanced binding to the second assay plate. Usually, the capture agent will be incubated in the wells for between about 8 to 72 hours to enable a sufficient coating of the capture agent to form on the inside walls of the wells. This incubation is generally carried out at low temperatures (e.g., between about 3-8°C) to avoid or reduce degradation of the capture agent.

Following incubation, the wells of the plate are decanted and tamped lightly with an absorbent substrate. Non-specific binding is then blocked. In order to achieve this, a block buffer, is added to the wells. For example, a block buffer containing bovine serum albumin (BSA) such as that sold by Intergen Company, Purchase, NY, is suitable. It has been found that 15 addition of between about 100 to  $200\mu$ l of block buffer to each well followed by gentle agitation at room temperature for between about 1-2 hours is sufficient to block non-specific binding. It is also possible to add the block buffer directly to the cell lysate obtained in the previous step rather than to the second assay plate.

Following this, the capture agent-coated plates are washed several times (usually between about 3-8 times) with a wash buffer. buffer can comprise phosphate buffered saline (PBS) at pH 7.0 to 7.5, for example. However, other wash buffers are available which can also be used. Conveniently, an automated plate washer, such as the ScanWasher 300 (Skatron Instruments, Inc., Sterling, VA) can be used for this, and other, washing steps of the assay.

## B. Measuring tyrosine phosphorylation

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The activated, solubilized rPTK (or receptor construct) is then added to the wells having the capture agent adhering thereto. As a general 30 proposition, about 80% of cell lysate obtained as mentioned under Section 1B above can be added to each well (i.e., about 60 to 160  $\mu$ l depending on the original volume of the wells). The lysate is incubated with the capture agent for an adequate period of time to enable the rPTK to be captured in the wells, e.g., from 1 to 3 hours. Incubation can be carried 35 out at room temperature.

Unbound cell lysate is then removed by washing with wash buffer. Following this washing step, an amount of the anti-phosphotyrosine antibody which is equal to, or less than, the amount of block buffer added 10

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previously, is added to each well. For example, about 50 to 200µl of an anti-phosphotyrosine antibody preparation having between about 0.3 to 0.5 µg/ml of antibody in a suitable buffer (e.g., PBS with a detergent such as those included in the lysis buffer) is added to the well. This is followed by a washing step to remove unbound anti-phosphotyrosine antibody.

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Tyrosine phosphorylation is then quantified by the amount of antiphosphotyrosine antibody binding to the second solid phase. Many systems
for detecting the presence of an antibody are available to those skilled
in the art. Some examples follow.

Generally, the anti-phosphotyrosine antibody will be labelled either directly or indirectly with a detectable label. Numerous labels are available which can be generally grouped into the following categories:

- (a) Radioisotopes, such as <sup>35</sup>S, <sup>14</sup>C, <sup>125</sup>I, <sup>3</sup>H, and <sup>131</sup>I. The antibody can be labeled with the radioisotope using the techniques described in Current Protocols in Immunology, supra, for example and radioactivity can be measured using scintillation counting.
  - (b) Fluorescent labels such as rare earth chelates (europium chelates) or fluorescein and its derivatives, rhodamine and its derivatives, dansyl, Lissamine, phycoerythrin and Texas Red are available. The fluorescent labels can be conjugated to the antibody using the techniques disclosed in <u>Current Protocols in Immunology</u>, supra, for example. Fluorescence can be quantified using a fluorimeter (Dynatech).
  - (c) Various enzyme-substrate labels are available and U.S. Patent No. 4,275,149 provides a review of some of these. The enzyme generally catalyses a chemical alteration of the chromogenic substrate which can be measured using various techniques. For example, the enzyme may catalyze a color change in a substrate, which can be measured spectrophotometrically. . Alternatively, the enzyme may alter the fluorescence or chemiluminescence of the substrate. Techniques for quantifying a change in fluorescence are described above. The chemiluminescent substrate becomes electronically excited by a chemical reaction and may then emit light which can be measured (using a Dynatech ML3000 chemiluminometer, for example) or donates energy to a fluorescent acceptor. Examples of enzymatic labels include luciferases (e.g., firefly luciferase and bacterial luciferase; U.S. Patent 2,3-dihydrophthalazinediones, 4,737,456), luciferin, dehydrogenase, urease, peroxidase such as horseradish peroxidase (HRPO), alkaline phosphatase, \$\beta\$-galactosidase, glucoamylase, lysozyme, saccharide oxidases (e.g., glucose oxidase, galactose oxidase, and glucose-6-phosphate

'dehydrogenase), heterocyclic oxidases (such as uricase and xanthine oxidase), lactoperoxidase, microperoxidase, and the like. Techniques for conjugating enzymes to antibodies are described in O'Sullivan et al., Methods for the Preparation of Enzyme-Antibody Conjugates for use in Enzyme Immunoassay, in Methods in Enzym. (ed J. Langone & H. Van Vunakis), Academic press, New York, 73: 147-166 (1981) and Current Protocols in Immunology, supra.

Examples of enzyme-substrate combinations include, for example:

- (i) Horseradish peroxidase (HRPO) with hydrogen peroxidase as a substrate, wherein the hydrogen peroxidase oxidizes a dye precursor (e.g. orthophenylene diamine [OPD] or 3,3',5,5'-tetramethyl benzidine hydrochloride [TMB]).
- (ii) alkaline phosphatase (AP) with para-Nitrophenyl phosphate as chromogenic substrate.
- (iii)  $\beta$ -D-galactosidase ( $\beta$ -D-Gal) with a chromogenic substrate (e.g. p-nitrophenyl- $\beta$ -D-galactosidase) or fluorogenic substrate 4-methylumbelliferyl- $\beta$ -D-galactosidase.

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Numerous other enzyme-substrate combinations are available to those skilled in the art. For a general review of these, see U.S. Patent Nos. 4,275,149 and 4,318,980.

Sometimes, the label is indirectly conjugated with the antibody. The skilled artisan will be aware of various techniques for achieving this. For example, the antibody can be conjugated with biotin and any of the three broad categories of labels mentioned above can be conjugated with avidin, or vice versa. Biotin binds selectively to avidin and thus, the label can be conjugated with the antibody in this indirect manner. See, Current Protocols in Immunology, supra, for a review of techniques involving biotin-avidin conjugation. Alternatively, to achieve indirect conjugation of the label with the antibody, the antibody is conjugated with a small hapten (e.g. digoxin) and one of the different types of labels mentioned above is conjugated with an anti-hapten antibody (e.g. anti-digoxin antibody). Thus, indirect conjugation of the label with the antibody can be achieved.

In another embodiment of the invention, the anti-phosphotyrosine antibody need not be labeled, and the presence thereof can be detected using a labeled anti-antiphosphotyrosine antibody (e.g. anti-mouse anti-phosphotyrosine antibody conjugated with HRPO).

In the preferred embodiment, the anti-phosphotyrosine antibody is labeled with an enzymatic label which catalyzes a color change of a substrate (such as tetramethyl benzimidine [TMB], or orthaphenylene diamine [OPD]). Thus, the use of radioactive materials is avoided. A color change of the reagent can be determined spectrophotometrically at a suitable wavelength (e.g. 450nm for TMB and 490nm for OPD, with a reference wavelength of 650 nm).

## 3. Intracellular Kinase Activity

The assay described herein is also useful for measuring phosphorylation and/or activation of intracellular kinases (e.g. cytoplasmic tyrosine kinases and/or cytoplasmic serine-threonine kinases). Phosphorylation of these molecules can occur as a consequence of transphosphorylation of the intracellular kinase by a kinase receptor or "receptor complex" (which comprises one or more kinase receptors residing in a cell membrane). Examples of intracellular tyrosine kinases include insulin receptor substrate I (IRS-1), Shc, Ras and GRB2, for example. Antibodies to human Shc, human Ras and GRB2 can be obtained commercially from UBI, NY, which can be used as capture agents for these tyrosine kinases. Examples of intracellular serine-threonine kinases include MEK and MAPK.

In order to measure phosphorylation of these kinases, the procedure is essentially as described above except that a chimera of the intracellular kinase and the flag polypeptide is normally formed (i.e. a "kinase construct"). Alternatively, the cell has an endogenous intracellular kinase or is transformed with nucleic acid encoding an intracellular kinase of interest. Generally, a eukaryotic cell will be transformed with nucleic acid encoding a kinase construct. Upon expression of the nucleic acid, the kinase or kinase construct will reside intracellularly (i.e. in the cytoplasm). The cells comprising the kinase or kinase construct are subjected to the KIRA as discussed above. Exposure to the agonist may result in trans-phosphorylation of the intracellular kinase which can be quantified in the ELISA as elaborated above. The capture agent in the ELISA binds to either the intracellular kinase or to the flag polypeptide.

## 4. Serine-Threonine Kinase Activity

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This assay is further useful for measuring phosphorylation and/or activation of serine-threonine kinases. The term "serine-threonine kinase" refers to a kinase which phosphorylates a substrate which has at least one

phosphate accepting alcohol group. The serine-threonine kinase is usually a "receptor" insofar as it has a ligand-binding ECD, TM domain and ICD. The ICD usually comprises a catalytic kinase domain and generally has one or more phosphate accepting serine and/or threonine residues. Examples of intracellular serine-threonine kinases include MEK and MAPK. See section 3 above for a discussion as to measuring phosphorylation of intracellular serine-threonine kinases. Examples of serine-threonine kinase receptors include daf-1, activin type II receptor (ActR-II), activin type IIB receptor (ActR-IIB), TGF-β type II receptor (TβR-II), activin receptor-like kinase (ALK) -1, -2, -3, -4 and TGF-β type I receptor (TβR-1)/ALK-5. See ten Dijke et al., supra. The serine-threonine kinase assay is essentially the same as described above for tyrosine kinases, except that phosphorylation is quantified using anti-phosphoserine and/or antiphosphothreonine antibodies. Anti-phosphoserine and anti-phosphothreonine 15 monoclonal antibodies can be purchased from Sigma Immuno Chemicals, St Louis, MO, for example.

## 5. Phosphatase Activity

Phosphatase activity can similarly be measured using the assay described herein. Phosphatase enzymes are able to dephosphorylate phosphorylated tyrosine, serine and/or threonine residues (i.e. liberate inorganic phosphate from phosphoric esters of such amino acid residues). Generally the phosphatase enzyme is specific for either tyrosine residues or serine-threonine residues but sometimes can dephosphorylate tyrosine, serine and threonine residues. Sometimes "endogenous" phosphatase activity is measured and this refers to the activity of phosphatase enzyme(s) which exist in nature in a selected cell.

In order to quantify endogenous phosphatase activity, cells possessing at least one phosphatase are stimulated in the presence and absence of one or more phosphatase inhibitors. Examples of protein tyrosine phosphatase (PTPase) inhibitors include sodium orthovanadate and sodium molybdate (Sigma Chemical Co., St. Louis, MO). ICN Biochemicals supply okadaic acid which is a serine-threonine phosphatase inhibitor. As a general proposition, between about 1-10µM phosphatase inhibitor can be added to each well of the assay plate. In all other respects, the assay is performed essentially as discussed above. Thus, the ability of endogenous phosphatases to dephosphorylate a kinase in the selected cell can be quantified.

In the preferred embodiment, a phosphatase enzyme of interest can be studied. Examples of protein tyrosine phosphatases (PTPases) include PTP1B, PTPMEG, PTP1c, Yop51, VH1, cdc25, CD45, HLAR, PTP18, HPTPc and DPTP10D. See Zhang and Dixon, Adv. Enzym. 68: 1-36 (1994). Examples of protein serine-5 threonine phosphatases include PP1, PP2A, PP2B and PP2C. See Meth. Enzym., ed Hunter & Sefton, Academic press, New York, 201:389-398 (1991). These proteins can be purchased commercially or made using the recombinant techniques described herein. To measure phosphatase activity, the KIRA ELISA can be performed essentially as described above with the following 10 modifications. Following capture of the kinase or kinase construct (e.g. receptor construct) to the second solid phase and the washing step (to remove unbound cell lysate), the phosphatase of interest is added to the wells of the second assay plate and incubated with the adhering kinase or kinase construct. For example, between about 50-200  $\mu$ l of the phosphatase 15 in a suitable dilution buffer (see Meth. Enzym., ed Hunter & Sefton, Academic press, New York, 201:416-440 [1991]) can be added to each well. This is generally followed by gentle agitation at room temperature (or 37°C) for between about 30 min to 2 hours to allow the phosphatase to dephosphorylate the kinase. Following washing to remove the phosphatase, the decreased degree of phosphorylation of the kinase relative to the control (i.e. no phosphatase added) is quantified by ELISA as described earlier herein.

# 6. Kits

As a matter of convenience, the reagents can be provided in a kit,

i.e., a packaged combination of reagents, for combination with the analyte
in assaying the ability of the analyte to activate or prevent activation
of a rPTK of interest. The components of the kit will be provided in
predetermined ratios. Thus, a kit will comprise the specific second solid
phase for the assay as well as the anti-flag polypeptide capture agent
either packaged separately or captured to the second solid phase (e.g. a
microtiter plate). Usually, other reagents, such as the antiphosphotyrosine antibody labelled directly or indirectly with an enzymatic
label will also be provided in the kit. Where the detectable label is an
enzyme, the kit will include substrates and cofactors required by the
centure (e.g. a substrate precursor which provides the detectable
chromophore or fluorophore). In addition, other additives may be included
such as stabilizers, buffers (e.g. a block buffer and a lysis buffer) and
the like. Conveniently, the kit can also supply the homogeneous population

of cells which have been transformed with the receptor construct. The relative amounts of the various reagents may be varied widely to provide for concentrations in solution of the reagents which substantially optimize the sensitivity of the assay. Particularly, the reagents may be provided as dry powders, usually lyophilized, including excipients which on dissolution will provide a reagent solution having the appropriate concentration. The kit also suitably includes instructions for carrying out the KIRA ELISA.

# 7. Uses for the Assay

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This application provides two assays which are useful for reliable, sensitive and quantitative detection of kinase activation. The first assay can be used where a kinase receptor-specific capture antibody having the desired characteristics herein described is available or has been prepared. The second assay is a generic assay which enables activation of any kinase receptor to be measured via the use of a flag polypeptide and a capture agent which binds specificity thereto.

These assays are useful for identifying novel agonists/antagonists for a selected kinase receptor. Also, the assay provides a means for studying ligand-receptor interactions (i.e., mechanism studies). Also the presence of an endogenous receptor in a selected cell line can be quantified using the assay. The assays are further useful for identifying the presence of a ligand for a selected kinase receptor in a biological sample and, e.g., establishing whether a growth factor has been isolated following a purification procedure. It is desirable to have an assay for measuring the ability of these growth factors to activate their respective receptors.

The assay also has clinical applications for detecting the presence of a ligand for a selected rPTK (e.g. the insulin receptor) in a biological sample taken from a human and thus patients having elevated or depressed levels of the ligand can be identified. This is particularly desirable where elevated or depressed levels of the ligand cause a pathological condition. Accordingly, candidates for administration of the selected ligand (e.g. insulin) can be identified through this diagnostic method. It is possible, using the assay disclosed herein, to assay the pK of agonists or antagonists administered to a patient. This assay also facilitates the detection of shed receptor in a biological sample.

The assay is also useful to quantify phosphatase activity of endogenous phosphatases or, in the preferred embodiment, a phosphatase of

interest. This can be used for screening phosphatase inhibitors, for example.

Below are examples of specific embodiments for carrying out the present inventi n. The examples are offered for illustrative purposes 5 only, and are not intended to limit the scope of the present invention in any way.

All publications, patents and patent applications cited herein, whether supra or infra, are hereby incorporated by reference in their entirety.

EXAMPLE 1

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### KIRA ELISA of the HER2 Receptor

The assay system described in this example was developed to measure the extent of autophosphorylation as a result of the interactions between the HER2 receptor and its specific activator, heregulin (HRG). 15 overexpression of p185HER2 has been correlated with poor clinical outcome in a number of epithelial-derived cancers. Heregulin and its rodent homologue, neu differentiation factor (NDF), were originally purified based on their ability to stimulate the autophosphorylation of a 185 kDa protein in the breast carcinoma cell lines MCF-7 and MDA-453, respectively. In this 20 embodiment of the invention, the cell line expressing the tyrosine kinase receptor DNA (either endogenous or transformed) is adherent and there is an antibody (e.g. monoclonal or affinity purified polyclonal) capable of specifically binding the receptor such that it neither stimulates autophosphorylation in the absence of ligand nor suffers impaired binding due to the presence of bound ligand. Standard curve preparations and many samples may easily be run simultaneously in replicate and at several dilutions using this assay, readily allowing quantitation of ligand activity in a large number of unknown samples.

# (i) Capture agent preparation

Polyclonal anti-HER2 antibody was isolated from pooled immune sera from New Zealand White rabbits immunized with the extracellular domain of the HER2 molecule (Fendly et al., Journal of Biological Response Modifiers The rHER2 ECD specific antibodies were affinity 9:449-455 [1990]). purified using an FPLC (Pharmacia Biotech, Inc, Piscataway, NJ) with an 35 affinity column generated from rHER2 ECD conjugated to Avidgel F (Bioprobe International, Inc., Tustin, CA). The resulting purified antibody stock was 0.829 mg/ml in phosphate buffered saline (PBS), pH 7.4, and was stored as 0.5 ml aliquots at -20°C.

### (ii) Anti-phosphotyrosine antibody preparation

Monoclonal anti-phosphotyrosine, clone 4G10, was purchased from 5 Upstate Biologicals, Inc (UBI, Lake Placid, NY) and biotinylated using long-arm biotin-N-hydroxysuccinamide (Biotin-X-NHS, Research Organics, Cleveland, OH).

#### (iii) Ligand

The recombinant truncated form of Blheregulin (MW= 7.88 Kd) 10 corresponding to residues 177-244 (HRG\$1,77-244) was produced in E. coli and purified to homogeneity as described in Holmes et al., Science, 256: 1205-1210 (1992) and was stored at 4°C as an 89.7  $\mu$ M stock solution in 50 mM Tris/HCl, pH 7.5.

## (iv) Adherent Cells

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MCF-7 (ATCC-HTB 22), an adherent cell line isolated from a human breast adenocarcinoma, was obtained from American Type Culture Collection (ATCC, Rockville, MD). MCF-7 cells have been shown to produce measurable levels of surface p185HER2 by both FACS and ELISA analysis. The cells were maintained in 150 cm2 tissue culture flasks (Corning Inc, Corning, NY) and 20 utilized when at cell densities of 60 % to 75 % confluency. For the assay,  $2 \times 10^5$  cells were seeded per well in flat-bottom microtiter plates (Falcon 3072, Becton Dickinson Labware, Lincoln Park, NJ) cultured overnight at 37°C in 5 % CO2. Cells were grown in F12/DMEM 50:50 Gibco as a custom formulation (Gibco/BRL, Life Technologies, Grand Island, NY). The medium was supplemented with 10 % FBS (HyClone, Logan, Utah), 25 mM HEPES (Gibco) and 2 mM L-glutamine (Gibco).

## (v) KIRA ELISA

MCF-7 cells (2 x  $10^5$ ) in 100  $\mu$ l media were added to each well in a flat-bottom-96 well culture plate and cultured overnight at 37°C in 5% CO<sub>2</sub>. The following morning the well supernatants were decanted, and the plates were lightly tamped on a paper towel. 50  $\mu$ l of media containing either experimental samples or the recombinant HRG\$1177-244 standards (3000, 1000, 333, 111, 37, 12, 4, and 0 pM) was then added to each well. The cells were stimulated at 37°C for 30 min., the well supernatants were decanted, and 35 the plates were once again lightly tamped on a paper towel. To lyse the cells and solubilize the receptors, 100  $\mu$ l of lysis buffer was added to each well. Lysis buffer consisted of 150 mM NaCl containing 50 mM HEPES (Gibco), 0.5 % Triton-X 100 (Gibco), 0.01 % thimerosal, 30 KIU/ml aprotinin (ICN Biochemicals, Aurora, OH), 1mM 4-(2-aminoethyl)-benzenesulfonyl fluoride hydrochloride (AEBSF; ICN Biochemicals), 50 μM leupeptin (ICN Biochemicals), and 2 mM sodium orthovanadate (Na<sub>3</sub>VO<sub>4</sub>, Sigma Chemical Co. St. Louis, MO), pH 7.5. The plate was then agitated gently on a plate shaker (Bellco Instruments, Vineland, NJ) for 60 min. at room temperature.

While the cells were being solubilized, an ELISA microtiter plate (Nunc Maxisorp. Inter Med, Denmark) coated overnight at 4°C with the affinity-purified polyclonal anti-HER2 ECD (1.0  $\mu$ g/ml in 50 mM carbonate buffer, pH 9.6, 100  $\mu$ l/well) was decanted, tamped on a paper towel and blocked with 150  $\mu$ l/well of Block Buffer [PBS containing 0.5 % BSA (Intergen Company, Purchase. NY) and 0.01 % thimerosal] for 60 min. at room temperature with gentle agitation. After 60 minutes, the anti-HER2 ECD coated plate was washed 6 times with wash buffer (PBS containing 0.05 % Tween-20 and 0.01 % thimerosal) using an automated plate washer (ScanWasher 300, Skatron Instruments, Inc. Sterling, VA).

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The lysate containing solubilized p185 HER2 from the cell-culture microtiter well was transferred (85  $\mu$ l/well) to anti-rHER2 ECD coated and blocked ELISA well and was incubated for 2 h at room temperature with gentle agitation. The unbound receptor was removed by washing with wash buffer and 100  $\mu$ l of biotinylated 4G10 (anti-phosphotyrosine) diluted 1:2000 in dilution buffer (PBS containing 0.5 % BSA, 0.05 % Tween-20, 5 mM EDTA, and 0.01 % thimerosal), i.e. 400pg/ml, was added to each well. After incubation for 2 h at room temperature the plate was washed and 100  $\mu$ l of HRPO-conjugated streptavidin (Zymed Laboratories, S. San Francisco, CA) 25 diluted 1:10000 in dilution buffer was added to each well. The plate was incubated for 30 minutes at room temperature with gentle agitation. The free avidin-conjugate was washed away and 100  $\mu$ l freshly prepared substrate solution (tetramethyl benzidine [TMB]; 2-component substrate kit; Kirkegaard and Perry, Gaithersburg, MD) was added to each well. reaction was allowed to proceed for 10 minutes, after which the color development was stopped by the addition of 100  $\mu$ l/well 1.0 M H<sub>1</sub>PO<sub>4</sub>. The absorbance at 450 nm was read with a reference wavelength of 650 nm (ABS<sub>450/650</sub>), using a vmax plate reader (Molecular Devices, Palo Alto, CA) controlled with a Macintosh Centris 650 (Apple Computers, Cupertino, CA) 35 and DeltaSoft software (BioMetallics, Inc, Princeton, NJ).

The standard curve shown in Figure 7 was generated by stimulating MCF-7 cells with 3000, 1000, 333, 111, 37, 12, 4, or 0 pM  $HRG\beta1_{177-244}$  and presented as pM  $HRG\beta1_{177-244}$  vs. mean  $ABS_{450/650}$   $\pm$  sd using the DeltaSoft

program. Sample concentrations were obtained by interpolation of their absorbance on the standard curve and are expressed in terms of pM  $HRG\beta1_{177}$ .

244 activity.

When the data were fitted to a 4-parameter nonlinear least squares equation, they resulted in a correlation coefficient of 0.9998. For the data shown in Figure 7, the EC<sub>50</sub> of receptor activation by  $HRG\beta1_{177-244}$  was 373 pM. To demonstrate the highly reproducible nature of the p185 KIRA ELISA, seven standard curves were generated over the period of one month and the EC<sub>50</sub>'s are averaged. This gives an EC<sub>50</sub>ave for  $HRG\beta1_{177-244}$  of 360  $\pm$  40 pM (average  $\pm$  SD).

### (vi) Intra- and inter-assay precision and assay specificity

The intra-assay variability was determined by performing the p185HER2 KIRA ELISA on three separate days. For each test, the standard curve is run in triplicate. Controls with HRG\$1177-244 corresponding to high (1000 pM), mid (200 pM) and low (40 pM) were assayed in 24 replicates. The ABS450/650 of the individual test samples were converted to pM HRG\$1177-244 activity and the 24 converted values for each test concentration were averaged. The data are expressed as averaged value and \* coefficient of variation (\*cv; [(intra-assay standard deviation/intra-assay averaged calculated value) x 100]. See Table 1A below.

Table 1

Intra- and Inter-assay Variation

A. Intra-assay Precision (n-24 per test)

High Value		Mid Value		Low Value	
Average Value (pM)	* cv <sup>b</sup>	Average Value (pM)	% cv	Average Value (pM)	% cv
1256 1078	19.5% 10.0%	209 196	10.8	33	12.3%
999	14.3%	196	5.1% 6.3%	38 35	7.5% 11.3%

Test #1 25 Test #2 Test #3

## B. Inter-assay Precision (n=3)

Average Value (pM)	₹cv <sup>c</sup>	Average Value (pM)	* cv	Average Value (pM)	* cv
1100	4.3%	200	6.3%	34	9.0%

Expected high value: 1000 pM; mid value: 200 pM; low value: 40 pM

b Intra-assay t cv determined as intra-assay sd/intra-assay average x 100

C Inter-assay t cv determined as inter-assay sd/inter-assay average x 100

The intra-assay variability of the KIRA ELISA was within acceptable limits despite the fact that the assay actually consists of both bioassay and ELISA components. The coefficients of variance (%) for the highest values were under 20% and for the mid and low values were at or under 10%.

The inter-assay variability was determined by averaging the values from upper-most three adjacent wells (of the 24 wells run) for a given sample concentration from each run. The three separate averages for each test concentration were then averaged. The data were expressed as averaged value and \*cv [(inter-assay standard deviation/inter-assay averaged value) x 100]. See Table 1B. above. The inter-assay variability of the KIRA ELISA was within acceptable limits.

In order to confirm the specificity of the assay, MCF-7 cells were stimulated with either HRG\$1<sub>177-244</sub> at 3000, 1000, 333, 111, 37, 12, 4 or 0 pM or insulin-like growth factor-1 (IGF-1), epidermal growth factor (EGF), vascular endothelial growth factor (VEGF), or insulin at 30000, 10000, 3333, 1111, 370, 120, 40 or 0 pM. The p185<sup>HER2</sup> KIRA ELISA was then performed as described above. The results are depicted in Figure 8.

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The p185<sup>HER2</sup> KIRA ELISA was clearly specific for heregulin. While HRGB1<sub>177-244</sub> induced normal receptor stimulation and autophosphorylation, the closely related EGF gives only a slight stimulation (OD<sub>450/650</sub> = 0.239) at the highest concentration tested (100 nM). Since EGF-R is produced in MCF-7 cells, this signal is likely due to EGF receptor transphosphorylation of p185<sup>HER2</sup>. Neither insulin-like growth factor-1 (IGF-1), vascular endothelial growth factor (VEGF) nor insulin have any detectable effect on the MCF-7 p185<sup>HER2</sup> KIRA ELISA, the latter despite the fact that MCF-7 cells produce active insulin receptors.

The results presented in this example demonstrate that the KIRA ELISA is a useful method for assaying ligand activation of a kinase receptor, e.g., heregulin activation of the  $p185^{HER2}$  receptor. Levels of receptor activation in terms of tyrosine phosphorylation are easily quantified and an EC<sub>50</sub> for a given ligand is readily determined. One potential use for this

assay would be to screen compounds for receptor agonist or antagonist activities. The potential throughput for this assay greatly surpasses that of Western blot analysis. Since the cell-culture portion of the assay is conducted in 96-well plates, many samples may be run in replicate at different dilutions at one time in a one-day assay.

### EXAMPLE 2

### KIRA ELISA of the Rse Receptor

Mark et al., Journal of Biological Chemistry 269(14):10720-10728 (1994) describe isolation of the Rse receptor protein tyrosine kinase from human and murine tissues. This Rse receptor with a carboxyl-terminal flag polypeptide (i.e. Rse.gD) was subjected to the KIRA ELISA described herein. The experimental procedure is outlined below.

### (i) Capture agent preparation

Monoclonal anti-gD (clone 5B6) was produced against a peptide from Herpes simplex virus glycoprotein D (Paborsky et al., <u>Protein Engineering</u> 3(6):547-553 [1990]). The purified stock preparation was adjusted to 3.0mg/ml in phosphate buffered saline (PBS), pH 7.4 and 1.0ml aliquots were stored at -20°C.

### (ii) Anti-phosphotyrosine antibody preparation

Monoclonal anti-phosphotyrosine, clone 4G10, was purchased from Upstate Biologicals, Inc (UBI, Lake Placid, NY) and biotinylated using long-arm biotin-N-hydroxysuccinamide (Biotin-X-NHS, Research Organics, Cleveland, OH).

## (iii) Ligand

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Since the endogenous ligand for the Rse receptor was not available, an agonist antibody for the Rse receptor was prepared which forms the ligand for the KIRA ELISA described in this Example. To generate the agonist antibody, a Rse.IgG chimera was generated. Briefly, the coding sequence of the ECD of Rse was fused to that of the human IgG-y1 heavy chain in a multi-step process. PCR was used to generate a fragment with a unique BstEII site 3' to the coding sequences of the Rse amino acid 428. The PCR product was joined to the human IgG-y1 heavy chain cDNA through a unique BstEII site in that construct (Mark et al., J. Cell. Biol., 267: 26166-26171 [1992]). The resulting construct (termed pRK.bpTK3.IgG.fusion) contained the coding sequences for amino acids 375-428 of Rse joined to those encoding human IgG-y1 heavy chain. The remaining portion of the Rse

ECD (amino acids 1-374) was then added by linkage through the Bam HI site in pRK.bpTK3.IgG.fusion to yield pRK.Rse.IgG.

To generate stable cell populations expressing Rse.IgG, the cDNA encoding Rse. IgG was subcloned into the episomal CMV-driven expression 5 plasmid pCIS.EBON, a pRK5 derivative disclosed in Cachianes et al., Bio. Techniques, 15: 225-259 (1993). Human fetal kidney 293 cells (obtained from ATCC, 12301 Parklawn Drive, Rockville, MD, USA) were transfected by the calcium phosphate technique. Cell monolayers were incubated for four hours in the presence of the DNA precipitate, glycerol shocked, and cultured in 10 F12:DMEM (1:1) containing 2mM glutamine, 10% fetal bovine serum, penicillin and streptomycin. After 48 hours, populations were replated in media containing G418 to select for a stable population of cells. Conditioned media was collected from cells expressing Rse. IgG nucleic acid that have been cultured in serum-free media for 72 hours in the absence of G418.

Rse. IqG was purified by affinity chromatography on a protein A column using procedures as described by Chamow, S.M., et al., Biochemistry, 29:9885-9891 (1990) with the following minor modifications. Conditioned media collected from cells expressing the Rse. IgG was adjusted to 0.1 M citrate pH 6.0 and loaded directly onto a protein A column (Repligen). The column was washed with 0.1 M citrate, pH 6.0, and was eluted with 3 M MgCl<sub>2</sub> with 10% glycerol. Fractions were pooled and desalted on a PD-10 column, dialyzed and concentrated against PBS. Protein concentrations were determined by an ELISA against human IgG (Fc). The protein was analyzed for purity by Coomassie staining of PAGE gels.

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Polyclonal antibodies were generated in New Zealand white rabbits against the Rse.IgG formed as described above.  $4\mu g$  of Rse.IgG in  $100\mu L$  PBS was emulsified with 100 L Freund's adjuvant (complete adjuvant for the primary injection and incomplete adjuvant for all boosts). For the primary immunization and the first boost, the protein was injected directly into 30 the popliteal lymph nodes (Sigel et al., Methods Enzymol., 93, 3-12 [1983]). For subsequent boosts, the protein was injected into subcutaneous and intramuscular sites. 1.3  $\mu$ g protein/kg body weight was injected every 3 weeks with bleeds taken 1 and 2 weeks following each boost. polyclonal antisera generated was then precipitated in 50% ammonium sulphate.

The resultant, purified polyclonal antisera is called "19B" herein. To confirm the ability of the 19B antisera to induce autophosphorylation of the Rse receptor, serum starved 3T3.gD.R11 cells (transformed with .

nucleic acid encoding the Rse receptor with an amino terminal gD flag polypeptide [i.e. gD.Rse] using the techniques described in Mark et al., Journal of Biological Chemistry 269(14):10720-10728 [1994]) or NIH3T3 cells were exposed to pre-immune serum or 19B polyclonal antisera at a 1:200 dilution for 10 minutes. The gD.Rse protein was immunoprecipitated from extracts using the anti-gD monoclonal antibody 5B6. Proteins were fractionated on 7% SDS-PAGE under reducing conditions and transferred to nitrocellulose. Phosphorylation of Rse was detected with labelled anti-phosphotyrosine antibody. Treatment of the 3T3.gD.R11 cells with 19B antisera stimulated the phosphorylation of the 140kD gD.Rse protein. This increase was not observed in cells treated with pre-immune sera.

The purified 19B polyclonal antisera was stored at 4°C as an 2.8mg/ml stock solution in PBS, pH 7.5.

### (iv) Preparation of Rse.qD nucleic acid

Synthetic double stranded oligonucleotides were used to reconstitute the coding sequence for the C-terminal 10 amino acids (880 - 890) of human Rse and add an additional 21 amino acids containing an epitope for the antibody 586 and a stop codon. The final sequence of the synthetic portion of the fusion gene was:

20 coding strand:

5'-TGCAGCAAGGGCTACTGCCACACTCGAGCTGCGCAGATGCTAGCCTCAAGATGGCT G
ATCCAAATCGATTCCGCGCCAAAGATCTTCCGGTCCTGTAGAAGCT-3' (SEQ ID NO: 10)
noncoding (anti-sense) strand:

5'-AGCTTCTACAGGACCGGAAGATCTTTGCCGCGGAATCGATTTGGATCAGCCATCTT G

25 AGGCTAGCATCTGCGCAGCTCGAGTGTGGCAGTAGCCCTTGCTGCA-3'(SEO ID NO: 11).

The synthetic DNA was ligated with the cDNA encoding amino acids 1-880 of human Rse at the PstI site beginning at nucleotide 2644 of the published human Rse cDNA sequence (Mark et al., Journal of Biological Chemistry 269(14):10720-10728 [1994]) and HindIII sites in the polylinker of the expression vector pSVI7.ID.LL (See Figure 16; SEQ ID NO: 9) to create the expression plasmid pSV.ID.Rse.gD. Briefly, the expression plasmid comprises a dicistronic primary transcript which contains sequence encoding DHFR bounded by 5' splice donor and 3' splice acceptor intron splice sites, followed by sequence that encodes the Rse.gD. The full length (non-spliced) message contains DHFR as the first open reading frame and therefore generates DHFR protein to allow selection of stable transformants.

# (v) Cell transformation

dp12.CHO cells (EP 307,247 published 15 March 1989) were electroporated with 20 μgs of pSV.ID.Rse.gD which had been linearized at a unique NotI site in the plasmid backbone. The DNA was ethanol precipitated after phenol/chloroform extraction and was resuspended in 20μl 1/10 Tris EDTA. Then, 10μg of DNA was incubated with 10<sup>7</sup> CHO.dp12 cells in 1 ml of PBS on ice for 10 min. before electroporation at 400 volts and 330μf. Cells were returned to ice for 10 min. before being plated into non-selective medium. After 24 hours cells were fed nucleoside-free medium to select for stable DHFR+ clones.

# (vi) Selection of transformed cells for use in the KIRA ELISA

To identify a cell line that expresses Rse.gD nucleic acid, candidate clones were screened by fluorescence activated cell sorting (FACS) analysis using the polyclonal antiserum 19B generated as described above, which recognizes epitopes in the extracellular domain of Rse. See Figure 5, step (b).

To confirm that clones that scored positive in the FACS assay express full-length Rse.gD nucleic acid, cell lysates were prepared (Lokker et al., EMBO J, 11:2503-2510 [1992]) and solubilized Rse.gD was immunoprecipitated 20 with the 19B antisera. The immunoprecipitated proteins were fractionated under reducing conditions using 7% PAGE, blotted onto nitrocellulose and then probed with the anti-gD 5B6 antibody which was detected with a horseradish peroxidase conjugated anti-mouse IgG antibody. See Figure 5, step (c). The ability of Rse.gD in cell clones to be activated to undergo autophosphorylation in response to the 19B agonistic antibody was determined. Briefly, serum starved dp.CHO cells transformed with Rse.gD nucleic acid as described above were exposed to pre-immune or 19B antisera at a 1:200 dilution for 10 min. The Rse.gD protein was immunoprecipitated from extracts using the anti-gD 5B6 monoclonal antibody. Proteins were 30 fractionated on 7% SDS-PAGE under reducing conditions and transferred to Phosphorylation of Rse was detected with labelled nitrocellulose. antiphosphotyrosine antibody. See Figure 5, step (d).

# (vii) Media

Cells were grown in F12/DMEM 50:50 (Gibco/BRL, Life Technologies, 35 Grand Island, NY). The media was supplemented with 10% diafiltered FBS (HyClone, Logan, Utah), 25mM HEPES and 2mM L-glutamine.

## (viii) KIRA ELISA

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Rse.gD transformed dp12.CHO cells (EP 307,247 published 15 March 1989) were seeded (5x104 per well) in the wells of a flat-bottom-96 well culture plate in 100  $\mu$ l media and cultured overnight at 37°C in 5% CO<sub>2</sub>. The following morning the well supernatants were decanted, and the plates were lightly tamped on a paper towel. 100µl of media containing either experimental samples or 1:100, 1:200, 1:400, 1:800, 1:1600, 1:3200 or 0 diluted, anti-Rse agonist polyclonal antibody (19B pAb) was then added to The cells were stimulated at 37°C for 30 min., the well supernatants were decanted, and the plates were once again lightly tamped on a paper towel. To lyse the cells and solubilize the receptors, 100  $\mu l$ of lysis buffer was added to each well. Lysis buffer consisted of 150 mM NaCl containing 50 mM HEPES (Gibco), 0.5 % Triton-X 100 (Gibco), 0.01 % thimerosal, 30 KIU/ml aprotinin (ICN Biochemicals, Aurora, OH), 1mM 4-(2hydrochloride ICN (AEBSF; 15 aminoethyl)-benzenesulfonyl fluoride Biochemicals), 50  $\mu$ M leupeptin (ICN Biochemicals), and 2 mM sodium orthovanadate (Na, VO, ; Sigma Chemical Co, St. Louis, MO), pH 7.5. The plate was then agitated gently on a plate shaker (Bellco Instruments, Vineland, NJ) for 60 min. at room temperature.

While the cells were being solubilized, an ELISA microtiter plate (Nunc Maxisorp, Inter Med, Denmark) coated overnight at 4°C with the 5B6 monoclonal anti-gD antibody (0.5  $\mu$ g/ml in 50 mM carbonate buffer, pH 9.6, 100  $\mu$ l/well) was decanted, tamped on a paper towel and blocked with 150  $\mu$ l/well of Block Buffer [PBS containing 0.5 % BSA (Intergen Company, Purchase, NY) and 0.01 % thimerosal] for 60 min. at room temperature with gentle agitation. After 60 minutes, the anti-gD 5B6 coated plate was washed 6 times with wash buffer (PBS containing 0.05 % Tween-20 and 0.01 % thimerosal) using an automated plate washer (ScanWasher 300, Skatron Instruments, Inc, Sterling, VA).

The lysate containing solubilized Rse.gD from the cell-culture microtiter well was transferred (85  $\mu$ l/well) to anti-gD 5B6 coated and blocked ELISA well and was incubated for 2 h at room temperature with gentle agitation. The unbound Rse.gD was removed by washing with wash buffer and 100  $\mu$ l of biotinylated 4G10 (anti-phosphotyrosine) diluted 1:2000 in dilution buffer (PBS containing 0.5 % BSA, 0.05 % Tween-20, 5 mM EDTA, and 0.01 % thimerosal), i.e. 400 pg/ml was added to each well. After incubation for 2 h at room temperature the plate was washed and 100  $\mu$ l of HRPO-conjugated streptavidin (Zymed Laboratories, S. San Francisco, CA)

diluted 1:10000 in dilution buffer was added to each well. The plate was incubated for 30 minutes at room temperature with gentle agitation. The free avidin-conjugate was washed away and 100 µl freshly prepared substrate solution (tetramethyl benzidine [TMB]; 2-component substrate kit; Kirkegaard and Perry, Gaithersburg, MD) was added to each well. The reaction was allowed to proceed for 10 minutes, after which the color development was stopped by the addition of 100 µl/well 1.0 M H<sub>3</sub>PO<sub>4</sub>. The absorbance at 450 nm was read with a reference wavelength of 650 nm (ABS<sub>450/650</sub>), using a vmax plate reader (Molecular Devices, Palo Alto, CA) controlled with a Macintosh Centris 650 (Apple Computers, Cupertino, CA) and DeltaSoft software (BioMetallics, Inc, Princeton, NJ).

The standard curve shown in Figure 10 was generated by stimulating Rse.gD transformed CHO cells with 1:100, 1:200, 1:400, 1:800, 1:1600, 1:3200 or 0 diluted, anti-Rse agonist antibody (19B) and presented as 1/dilution anti-Rse agonist antibody (19B) vs. mean  $ABS_{450/650} \pm \text{sd}$  using the DeltaSoft program.

The results presented in this example demonstrate that the KIRA ELISA is a useful method for assaying ligand activation of a receptor construct having a carboxyl terminal flag polypeptide, e.g., activation of Rse.gD. Levels of receptor activation in terms of tyrosine phosphorylation are easily quantified and an  $EC_{50}$  for a given ligand (e.g. an agonist antibody for the receptor) is readily determined.

## EXAMPLE 3

# KIRA ELISA of the trk A. B and C Receptors

Neurotrophins belong to a family of small, basic proteins which play a crucial role in the development and maintenance of the nervous system. The first identified and probably best understood member of this family is nerve growth factor (NGF). See U.S. Patent No. 5,169,762, issued December 8, 1992. Recently, sequentially related but distinct polypeptides with similar functions to NGF have been identified. For example, brain-derived neurotrophic factor (BDNF), now also referred to as neurotrophin-2 (NT2), was cloned and sequenced by Leibrock et al. (Nature, 341: 149-152 [1989]). Several groups identified a neurotrophic factor originally called neuronal factor (NF), and now referred to as neurotrophin-3 (NT3). (Ernfors et al., Proc. Natl. Acad. Sci. USA, 87: 5454-5458 [1990]; Köhn et al., Nature, 344: 339 [1990]; Maisonpierre et al., Science, 247: 1446 [1990]; Rosenthal et al., Neuron, 4: 767 [1990]; Jones and Reichardt, Proc. Natl. Acad. Sci.

USA, 87: 8060-8064 [1990]; Kaisho et al., FEBS\_Lett., 266: 187 [1990]).
Neurotrophins-4 and -5 (NT4 and NT5) have been recently added to the family
(Hallbook et al., Neuron, 6: 845-858 [1991]; Berkmeier et al., Neuron, 7:
857-866 [1991]; Ip et al., Proc. Natl. Acad. Sci. USA, 89: 3060-3064
5 [1992]).

Neurotrophins, similarly to other polypeptide growth factors, affect their target cells through interactions with cell surface rPTKs (called Trk receptors). The first member of the trk receptor family, trkA, was initially identified as the result of an oncogenic transformation caused 10 by the translocation of tropomyosin sequences onto its catalytic domain. Later work identified trkA as a signal transducing receptor for NGF. Subsequently, two other related receptors, mouse and rat trkB (Klein et al., EMBO J., 8: 3701-3709 [1989]; Middlemas et al., Mol. Cell. Biol., 11: 143-153 [1991]; EP 455,460 published 6 November 1991) and porcine, mouse and rat trkC (Lamballe et al., Cell, 66: 967-979 [1991]; EP 522,530 published 13 January 1993), were identified as members of the trk receptor family. The structures of the trk receptors are quite similar, but alternate splicing increases the complexity of the family by giving rise to two known forms of trkA, three known forms of trkB (two without 20 functional tyrosine kinase domains) and at least four forms of trkC (several without functional tyrosine kinase domain, and two with small inserts in the tyrosine kinase domain). Human trk A, B and C receptor sequences are disclosed in U.S. Patent application Serial No. 08/215,139, filed March 18, 1994, specifically incorporated herein by reference.

The following KIRA ELISA was performed using trk A, B and C receptor constructs having amino-terminal flag polypeptides.

# (i) Capture agent preparation

Monoclonal anti-gD (clone 5B6) was produced against a peptide from Herpes simplex virus glycoprotein D as discussed above in Example 2. The purified stock preparation was adjusted to 3.0mg/ml in phosphate buffered saline (PBS), pH 7.4 and 1.0ml aliquots were stored at -20°C.

# (ii) Anti-phosphotyrosine antibody preparation

Monoclonal anti-phosphotyrosine, clone 4G10, was purchased from Upstate Biologicals, Inc (UBI, Lake Placid, NY) and biotinylated using long-arm biotin-N-hydroxysuccinamide (Biotin-X-NHS, Research Organics, Cleveland, OH).

## (iii) Ligands

Nerve growth factor (NGF), neurotrophin 3 (NT3), and neurotrophin 5 (NT5) were prepared by recombinant techniques using the sequence data provided for each of these proteins in the above-mentioned references. The purified NGF, NT3 and NT5 were stored at 4°C as stock solutions (180 µM, 8.8 µM and 26.9 µM, respectively) in PBS, pH 7.5.

## (iv) Preparation of gD.trk nucleic acid

In order to express the various trk receptors with gD flags (i.e. gD.trk constructs), DNA constructs were made which encoded the signal and epitope of gD (see Paborsky et al., supra) fused to the amino terminus of 10 the various trk receptors. These were made by inserting the trk receptor and gD sequences into pRK5 or pRK7 (Suva et al., Science, 237: 893-896 [1987]) using standard molecular biology techniques, to generate the constructs shown in Figures 12-14. In addition to the gD.trk constructs, constructs were also made to express gD tagged trk. IgG fusion proteins DNA constructs encoding the chimeras of trk 15 (i.e., gD.trk.IgG). extracellular domain and IgG-1 Fc domains were made with the Pc region clones of human IgG-1 (Ashkenazi et al., Immunoadhesins Intern. Rev. Immunol., 10: 219-227 [1993]). More specifically, the source of the IgG-1 encoding sequence was the CD4-IgG-1 expression plasmid pRKCD42Fc1 (Capon et al., Nature, 334: 525 [1989]; Byrn et al., Nature, 344: 667 [1990]) containing a cDNA sequence encoding a hybrid polypeptide consisting of residues 1-180 of the mature human CD4 protein fused to human IgG-1 sequences beginning at aspartic acid 216 (taking amino acid 114 as the first residue of the heavy chain constant region; Kabat et al., Sequences of Proteins of Immunological Interest 4th ed. [1987]), which is the first residue of the IgG-1 hinge after the cysteine residue involved in heavylight chain bonding, and ending with residues 441 to include the CH2 and CH3 Fc domains of IgG-1. The CD4-encoding sequence was deleted from the expression plasmid pRKCD42Fc1 and the vector was fused to DNA encoding the trk receptors, with the splice between aspartate 216 of the IgG-1 and valine 402 of trkA, threonine 422 of trkB, or threonine 413 of trkC. The gD tag was added to the amino terminus of each trk. IgG in the same way as for the gD.trk constructs.

# (v) Cell transformation

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Human embryonic kidney 293 cells (obtained from ATCC, Rockville, MD) were transiently transfected with the nucleic acid encoding gD.trk.IgG using a calcium phosphate protocol (Gorman, DNA\_Cloning: A Practical Approach [Glover, D., ed.] Vol II: 143-190, IRL Press, Washington DC).

After twelve hours, the transformed cells were rinsed three times with serum free F12/DMEM 50:50 media (Gibco) and then serum free media was added for a 48 hour collection.

cell lines stably expressing each of the gD.trk constructs were made by co-transfecting dp12.CHO cells (EP 307.247 published 15 March 1989) with the pRK plasmids encoding the gD tagged trk receptors and a plasmid encoding DHFR, again using calcium phosphate mediated transfection.

The media mentioned above (having the gD.trk.IgG) was used without further purification in binding assays to assess the effects of the presence of the gD flag polypeptide on neurotrophin binding to the gD.trk.IgG polypeptides. DNA encoding untagged trk.IgG polypeptide was run in parallel as a control. trk.IgG and gD tagged trk.IgG containing cell supernatants were prepared as described and used in competitive displacement assays with the appropriate iodinated neurotrophin. NGF is used as ligand for trkA, NT5 is used as ligand for trkB, and NT3 is used as a ligand for trkC. A summary of the results obtained is shown in the following table.

TABLE 1
Binding of Neurotrophins to trk.IgG

		IC50 without gD	IC50 with gD
20	trkA	68.4+/-11.9 pM	68.8+/-3.0 pM
	trkB	31.1+/-15.6 pM	12.1+/-18 pM
	trkC	31.1+/-1.1 pM	30.2+/-0.7 pM

# (vi) Selection of transformed cells for use in the KIRA ELISA

It was apparent from the preceding experiment that there was no observable change in the affinity of interaction of neurotrophins with their receptor due to the presence of the gD flag polypeptide on the amino terminus. Based on this result, cells were transformed with the gD.trk constructs for use in the KIRA ELISA using the techniques described in the previous section.

After two days, dp12.CHO cells (EP 307,247 published 15 March 1989) transformed with gD.trk constructs were selected for by growth in media without GHT, and after two weeks, growing cells were sorted by FACS analysis using the 5B6 monoclonal to select cells expressing the gD flag polypeptide on their surface. gD positive cells were cloned by plating at limiting dilution and resultant colonies were then rescreened by FACS

analysis (using the anti-gD 5B6 monoclonal antibody), neurotrophin binding (as discussed above), tyrosine phosphorylation indicated by Western blot using an anti-phosphotyrosine antibody, gD expression by Western blot using th anti-gD 5B6 antibody, and immunocytochemistry using the 5B6 antibody. 5 Clones which were positive were then recloned by limiting dilution and were subjected to the KIRA ELISA as described below.

## (vii) Media

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Cells were grown in F12/DMEM 50:50 (Gibco/BRL, Life Technologies, Grand Island, NY). The media was supplemented with 10% diafiltered FBS (HyClone, Logan, Utah), 25mM HEPES and 2mM L-glutamine.

## (viii) KIRA ELISA

gD.trk transformed dp12.CHO cells (EP 307,247 published 15 March 1989) were seeded (5 x 104 per well) in a flat-bottom-96 well culture plate in 100  $\mu$ l media and cultured overnight at 37°C in 5% CO<sub>2</sub>. The following morning the well supernatants were decanted, and the plates were lightly tamped on a paper towel.  $100\mu$ l of media containing either experimental samples or the recombinant purified NGF, NT3, or NT5 standards (3000, 1000, 333, 111, 37, 12, 4, and 0 pM) was then added to each well. The cells were stimulated at 37°C for 30 min., the well supernatants were decanted, and the plates were once again lightly tamped on a paper towel. To lyse the cells and solubilize the receptors, 100 µl of lysis buffer was added to each well. Lysis buffer consisted of 150 mM NaCl containing 50 mM HEPES (Gibco), 0.5 % Triton-X 100 (Gibco), 0.01 % thimerosal, 30 KIU/ml aprotinin (ICN Biochemicals, Aurora, OH), 1mM 4-(2-aminoethyl)-benzenesulfonyl fluoride hydrochloride (AEBSF; ICN Biochemicals), 50 µM leupeptin (ICN Biochemicals), and 2 mM sodium orthovanadate (Na<sub>3</sub>VO<sub>4</sub>; Sigma Chemical Co, St. Louis, MO). pH 7.5. The plate was then agitated gently on a plate shaker (Bellco Instruments, Vineland, NJ) for 60 min. at room temperature.

While the cells were being solubilized, an ELISA microtiter plate (Nunc Maxisorp, Inter Med, Denmark) coated overnight at 4°C with the 5B6 monoclonal anti-gD antibody (0.5  $\mu$ g/ml in 50 mM carbonate buffer, pH 9.6, 100  $\mu$ l/well) was decanted, tamped on a paper towel and blocked with 150  $\mu$ l/well of Block Buffer (PBS containing 0.5 % BSA (Intergen Company, Purchase, NY) and 0.01 % thimerosal] for 60 min. at room temperature with 35 gentle agitation. After 60 minutes, the anti-gD 5B6 coated plate was washed 6 times with wash buffer (PBS containing 0.05 % Tween-20 and 0.01 % thimerosal) using an automated plate washer (ScanWasher 300, Skatron Instruments, Inc. Sterling, VA).

The lysate containing solubilized gD.trk from the cell-culture microtiter well was transferred (85  $\mu$ l/well) to anti-gD 5B6 coated and blocked ELISA well and was incubated for 2 h at room temperature with gentle agitation. The unbound gD.trk was removed by washing with wash 5 buffer and 100  $\mu$ l of biotinylated 4G10 (anti-phosphotyrosine) diluted 1:2000 in dilution buffer (PBS containing 0.5 % BSA, 0.05 % Tween-20, 5 mM EDTA, and 0.01 % thimerosal), i.e., 400pg/ml, was added to each well. After incubation for 2 h at room temperature the plate was washed and 100 ul of HRPO-conjugated streptavidin (Zymed Laboratories, S. San Francisco, CA) diluted 1:10000 in dilution buffer was added to each well. The plate was incubated for 30 minutes at room temperature with gentle agitation. The free avidin-conjugate was washed away and 100  $\mu$ l freshly prepared substrate solution (tetramethyl benzidine; 2-component substrate kit; Kirkegaard and Perry, Gaithersburg, MD) was added to each well. 15 reaction was allowed to proceed for 10 minutes, after which the color development was stopped by the addition of 100  $\mu$ l/well 1.0 M H<sub>3</sub>PO<sub>4</sub>. The absorbance at 450 nm was read with a reference wavelength of 650 nm (ABS<sub>450/650</sub>), using a vmax plate reader (Molecular Devices, Palo Alto, CA) controlled with a Macintosh Centris 650 (Apple Computers, Cupertino, CA) 20 and DeltaSoft software (BioMetallics, Inc, Princeton, NJ).

The standard curves shown in Figures 15A-15C were generated by stimulating gD.trk transformed CHO cells with 3000, 1000, 333, 111, 37, 12, 4, and 0 pM NGF, NT3 or NT5 and were presented as pM neurotrophin vs. mean ABS<sub>450/650</sub> ± sd using the DeltaSoft program. Sample concentrations were obtained by interpolation of their absorbance on the standard curve and are expressed in terms of pM neurotrophin activity.

The results presented in this example demonstrate that the KIRA ELISA is a useful method for assaying ligand activation of a receptor construct having an amino terminal flag polypeptide, e.g., activation of gD.trk receptor constructs. Levels of receptor activation in terms of tyrosine phosphorylation are easily quantified and an EC50 for a given ligand is readily determined.

## EXAMPLE 4

## KIRA ELISA of the MPL/Rse Chimeric Receptor

The human MPL receptor has been disclosed by Vigon et al., <u>PNAS. USA</u>
89:5640-5644 (1992). A chimeric receptor comprising the ECD of the MPL receptor and the TM and ICD of Rse (Mark et al., supra) with a carboxyl-

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terminal flag polypeptide (i.e. Rse.gD; see Example 2) was subjected to the KIRA ELISA described herein. The experimental procedure is outlined below. See also Figs. 16 and 17.

# (i) Capture agent preparation

Monoclonal anti-gD (clone 5B6) was produced against a peptide from Herpes simplex virus glycoprotein D (Paborsky et al., Protein Engineering 3(6):547-553 [1990]). The purified stock preparation was adjusted to 3.0mg/ml in phosphate buffered saline (PBS), pH 7.4 and 1.0ml aliquots were stored at -20°C.

# (ii) Anti-phosphotyrosine antibody preparation

Monoclonal anti-phosphotyrosine, clone 4G10, was purchased from UBI (Lake Placid, NY) and biotinylated using long-arm biotin-N-hydroxysuccinamide (Biotin-X-NHS, Research Organics, Cleveland, OH).

### (iii) Ligand

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The MPL ligand [de Sauvage et al., Nature 369: 533-538 (1994)] was prepared by recombinant techniques. The purified MPL ligand was stored at 4°C as a stock solution.

# (iv) Preparation of MPL/Rse.gD nucleic acid

The expression plasmid pSV.ID.Rse.gD produced as described in Example 20 2 above was modified to produce plasmid pSV.ID.M.tmRd6 which contained the coding sequences of the ECD of human MPL (amino acids 1-491) fused to the transmembrane domain and intracellular domain of Rse.gD (amino acids 429-911). Synthetic oligonucleotides were used to join the coding sequence of a portion of the extracellular domain of human MPL to a portion of the Rse coding sequence in a two step PCR cloning reaction as described by Mark et al. in J. Biol. Chem. 267: 26166-26171 (1992). Primers used for the first PCR reaction were M1 (5'-TCTCGCTACCGTTTACAG - SEQ ID NO:12) and M2 (5'-CAGGTACCCACCAGGCGGTCTCGGT - SEQ ID NO: 13) with a MPL cDNA template and R1 (5'-14) R2 (5'-GGGCCATGACACTGTCAA SEQ NO: and 30 GACCGCCACCGAGACCGCCTGGTGGGTACCTGTGGTCCTT - SEQ ID NO: 15) with a Rse cDNA template. The PvuII-SmaI portion of this fusion junction was used for the construction of the full-length chimeric receptor.

# (v) Cell transformation

dp12.CHO cells (EP 307,247 published 15 March 1989) were electroporated with pSV.ID.M.tmRd6 which had been linearized at a unique NotI site in the plasmid backbone. The DNA was ethanol precipitated after phenol/chloroform extraction and was resuspended in  $20\mu l$  1/10 Tris EDTA. Then,  $10\mu g$  of DNA was incubated with  $10^7$  CHO.dp12 cells in 1 ml of PBS on

returned to ice for 10 min. before being plated into non-selective medium.

After 24 hours cells were fed nucleoside-free medium to select for stable

DHFR+ clones.

5 (vi) Selection of transformed cells for use in the KIRA ELISA

Clones expressing MPL/Rse.gD were identified by western-blotting of whole cell lysates post-fractionation by SDS-PAGE using the antibody 5B6 which detects the gD epitope tag.

### (vii) Media

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Cells were grown in F12/DMEM 50:50 (Gibco/BRL, Life Technologies, Grand Island, NY). The media was supplemented with 10% diafiltered FBS (HyClone, Logan, Utah), 25mM HEPES and 2mM L-glutamine.

## (viii) KIRA ELISA

MPL/Rse.gD transformed dp12.CHO cells were seeded (3x104 per well) in 15 the wells of a flat-bottom-96 well culture plate in 100  $\mu$ l media and cultured overnight at 37°C in 5% CO2. The following morning the well supernatants were decanted, and the plates were lightly tamped on a paper towel.  $50\mu$ l of media containing either experimental samples or 200, 50, 12.5, 3.12, 0.78, 0.19, 0.048 or 0 ng/ml MPL ligand was then added to each 20 well. The cells were stimulated at 37°C for 30 min., the well supernatants were decanted, and the plates were once again lightly tamped on a paper towel. To lyse the cells and solubilize the chimeric receptors, 100  $\mu$ l of lysis buffer was added to each well. Lysis buffer consisted of 150 mM NaCl containing 50 mM HEPES (Gibco), 0.5 % Triton-X 100 (Gibco), 0.01 % 25 thimerosal, 30 KIU/ml aprotinin (ICN Biochemicals, Aurora, OH), 1mM 4-(2-Biochemicals), 50  $\mu M$  leupeptin (ICN Biochemicals), and 2 mM sodium orthovanadate (Na3VO4; Sigma Chemical Co, St. Louis, MO), pH 7.5. The plate was then agitated gently on a plate shaker (Bellco Instruments, Vineland, NJ) for 60 min. at room temperature.

While the cells were being solubilized, an ELISA microtiter plate (Nunc Maxisorp, Inter Med, Denmark) coated overnight at 4°C with the 5B6 monoclonal anti-gD antibody (5.0  $\mu$ g/ml in 50 mM carbonate buffer, pH 9.6, 100  $\mu$ l/well) was decanted, tamped on a paper towel and blocked with 150  $\mu$ l/well of Block Buffer [PBS containing 0.5 % BSA (Intergen Company, Purchase, NY) and 0.01 % thimerosal] for 60 min. at room temperature with gentle agitation. After 60 minutes, the anti-gD 5B6 coated plate was washed 6 times with wash buffer (PBS containing 0.05 % Tween-20 and 0.01 %

thimerosal) using an automated plate washer (ScanWasher 300, Skatron Instruments, Inc., Sterling, VA).

The lysate containing solubilized MPL/Rse.gD from the cell-culture microtiter well was transferred (85  $\mu$ l/well) to anti-gD 5B6 coated and 5 blocked ELISA well and was incubated for 2 h at room temperature with gentle agitation. The unbound MPL/Rse.gD was removed by washing with wash buffer and 100  $\mu$ l of biotinylated 4G10 (anti-phosphotyrosine) diluted 1:18000 in dilution buffer (PBS containing 0.5 % BSA, 0.05 % Tween-20, 5 mM EDTA, and 0.01 % thimerosal), i.e. 56 ng/ml was added to each well. 10 After incubation for 2 h at room temperature the plate was washed and 100 μl of HRPO-conjugated streptavidin (Zymed Laboratories, S. San Francisco, CA) diluted 1:60000 in dilution buffer was added to each well. The plate was incubated for 30 minutes at room temperature with gentle agitation. The free avidin-conjugate was washed away and 100  $\mu$ l freshly prepared 15 substrate solution (tetramethyl benzidine [TMB]; 2-component substrate kit; Kirkegaard and Perry, Gaithersburg, MD) was added to each well. reaction was allowed to proceed for 10 minutes, after which the color development was stopped by the addition of 100  $\mu$ l/well 1.0 M H<sub>3</sub>PO<sub>4</sub>. The absorbance at 450 nm was read with a reference wavelength of 650 nm 20 (ABS<sub>450/650</sub>), using a vmax plate reader (Molecular Devices, Palo Alto, CA) controlled with a Macintosh Centris 650 (Apple Computers, Cupertino, CA) and DeltaSoft software (BioMetallics, Inc, Princeton, NJ).

The results demonstrated that MPL ligand was able to activate the MPL/Rse.gD chimeric receptor in a concentration-dependent and ligandspecific manner.

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#### **CLAIMS**

- 1. A method for measuring autophosphorylation of a tyrosine kinase receptor comprising the steps of:
  - (a) coating a first solid phase with a homogeneous population of eukaryotic cells so that the cells adhere to the first solid phase, wherein, positioned in their membranes, the cells have a receptor construct comprising a flag polypeptide and the tyrosine kinase receptor;
  - (b) exposing the adhering cells to an analyte;
- (c) solubilizing the adhering cells, thereby releasing cell lysate therefrom;
  - (d) coating a second solid phase with a capture agent which binds specifically to the flag polypeptide so that the capture agent adheres to the second solid phase;
- (e) exposing the adhering capture agent to the cell lysate obtained in step (c) so that the receptor construct adheres to the second solid phase;
  - (f) washing the second solid phase so as to remove unbound cell lysate;
- 20 (g) exposing the adhering receptor construct to an antiphosphotyrosine antibody which identifies phosphorylated
  tyrosine residues in the tyrosine kinase receptor; and
  - (h) measuring binding of the anti-phosphotyrosine antibody to the adhering receptor construct.
- 25 2. The method of claim 1 wherein the cells are transformed with nucleic acid encoding the receptor construct prior to step (a).
  - 3. The method of claim 1 wherein the cells comprise a mammalian cell line.
  - 4. The method of claim 1 wherein the cells are adherent.
- 30 5. The method of claim 1 wherein the capture agent comprises a capture antibody.

- 6. The method of claim 1 wherein the first solid phase comprises a well of a first assay plate.
- 7. The method of claim 6 wherein the first assay plate is a microtiter plate.
- 5 8. The method of claim 6 wherein between about 1 x  $10^4$  to 3 x  $10^5$  cells are added to the well in step (a).
  - 9. The method of claim 1 wherein the second solid phase comprises a well of a second assay plate.
- 10. The method of claim 1 wherein the cell lysate is not concentrated or clarified prior to step (e).
  - 11. The method of claim 6 wherein step (c) comprises adding a lysis buffer to the well of the first assay plate and gently agitating the first assay plate.
- 12. The method of claim 11 wherein the lysis buffer comprises a solubilizing detergent.
  - 13. The method of claim 1 wherein the anti-phosphotyrosine antibody is labelled.
- 14. The method of claim 13 wherein the label comprises an enzyme which is exposed to a color reagent and the color change of the color reagent is determined in step (h).
  - 15. The method of claim 1 wherein the flag polypeptide is fused to the amino terminus of the tyrosine kinase receptor.
  - 16. The method of claim 15 wherein the tyrosine kinase receptor is a trk
    A receptor, trk B receptor or trk C receptor.
- 25 17. The method of claim 1 wherein the flag polypeptide is fused to the carboxyl terminus of the tyrosine kinase receptor.

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18. The method of claim 17 wherein the tyrosine kinase receptor is the Rse receptor.

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- 19. The method of claim 17 wherein the receptor construct comprises the extracellular domain of a receptor of interest and the intracellular domain of the Rse receptor.
  - 20. The method of claim 19 wherein the receptor of interest is the MPL receptor.
- 21. The method of claim 20 wherein the receptor construct further comprises the transmembrane domain of the Rse receptor and the flag polypeptide comprises the gD polypeptide.
  - 22. The method of claim 1 wherein the analyte comprises an agonist for the tyrosine kinase receptor.
- 23. The method of claim 1 wherein the analyte comprises an antagonist for the tyrosine kinase receptor.
- 15 24. The method of claim 23 wherein the antagonist competitively inhibits binding or activation of the tyrosine kinase receptor by an agonist thereto and step (b) is followed by a step wherein the adhering cells are exposed to the agonist.
- 25. The method of claim 1 wherein the analyte is a composition which comprises an antagonist and an agonist for the receptor and the assay measures the ability of the antagonist to bind to the agonist and thereby reduce activation of the tyrosine kinase receptor by the agonist.
- 26. The method of claim 1 wherein a block buffer is added to the second solid phase following step (d).
  - 27. A method for measuring autophosphorylation of a tyrosine kinase receptor comprising the steps of:
    - (a) coating a well of a first assay plate with a homogeneous population of adherent cells so that the cells adhere to the

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- well, wherein the cells have a tyrosine kinase receptor positioned in the cell membranes thereof;
- exposing the adhering cells to an analyte; (b)
- solubilizing the adhering cells thereby releasing cell lysate (c) therefrom;
- coating a well of a second assay plate with a capture agent (d) which binds specifically to the tyrosine kinase receptor so that the capture agent adheres to the well;
- (e) exposing the cell lysate obtained in step (c) to the adhering capture agent so that the tyrosine kinase receptor adheres to 10 the well;
  - washing the well so as to remove unbound cell lysate; (f)
  - exposing the adhering tyrosine kinase receptor to an anti-(g) phosphotyrosine antibody which binds selectively to phosphorylated tyrosine residues in the tyrosine kinase receptor;
  - measuring binding of the anti-phosphotyrosine antibody to the (h) adhering tyrosine kinase receptor.
- The method of claim 27 wherein the tyrosine kinase receptor comprises 28. the HER2 receptor. 20
  - A polypeptide comprising a flag polypeptide fused to the carboxyl 29. terminus of the intracellular domain of the Rse receptor.
  - The polypeptide of claim 29 further comprising the transmembrane domain of the Rse receptor.
- The polypeptide of claim 30 further comprising the extracellular 31. 25 domain of a receptor protein tyrosine kinase, other than the Rse receptor.
  - The polypeptide of claim 29 wherein the flag polypeptide comprises 32. the gD flag.
- A kit comprising a solid phase coated with a capture agent which 30 33. binds specifically to a flag polypeptide.

- 34. The kit of claim 33 wherein the solid phase comprises a well of a microtiter plate.
- 35. The kit of claim 33 further comprising a labeled anti-phosphotyrosine antibody.
- 5 36. The kit of claim 35 wherein the label comprises an enzyme.
  - 37. The kit of claim 33 further comprising a cell transformed with a nucleic acid encoding a receptor construct.
  - 38. An assay for measuring phosphorylation of a kinase comprising the steps of:
- 10 (a) coating a first solid phase with a homogeneous population of eukaryotic cells so that the cells adhere to the first solid phase, wherein the cells comprise a kinase construct comprising a flag polypeptide and the kinase;
  - (b) exposing the adhering cells to an analyte;
- 15 (c) solubilizing the adhering cells, thereby releasing cell lysate therefrom;
  - (d) coating a second solid phase with a capture agent which binds specifically to the flag polypeptide so that the capture agent adheres to the second solid phase;
- 20 (e) exposing the adhering capture agent to the cell lysate obtained in step (c) so that the kinase construct adheres to the second solid phase;
  - (f) washing the second solid phase so as to remove unbound cell lysate;
- 25 (g) exposing the adhering kinase construct to an antibody which identifies phosphorylated residues in the kinase construct; and
  - (h) measuring binding of the antibody to the adhering kinase construct.
  - 39. The assay of claim 38 wherein the kinase is a receptor.
- 30 40. The assay of claim 38 wherein the kinase is a serine-threonine kinase.

- 41. The assay of claim 38 which measures phosphatase activity.
- The assay of claim 41 wherein the cells further comprise a phosphatase and the assay further comprises the step of exposing the eukaryotic cells to a phosphatase inhibitor prior to step (c).
- 5 43. The assay of claim 41 which further comprises the steps in between steps (f) and (g) of exposing the adhering kinase construct to a phosphatase and then washing the second solid phase so as to remove unbound phosphatase.

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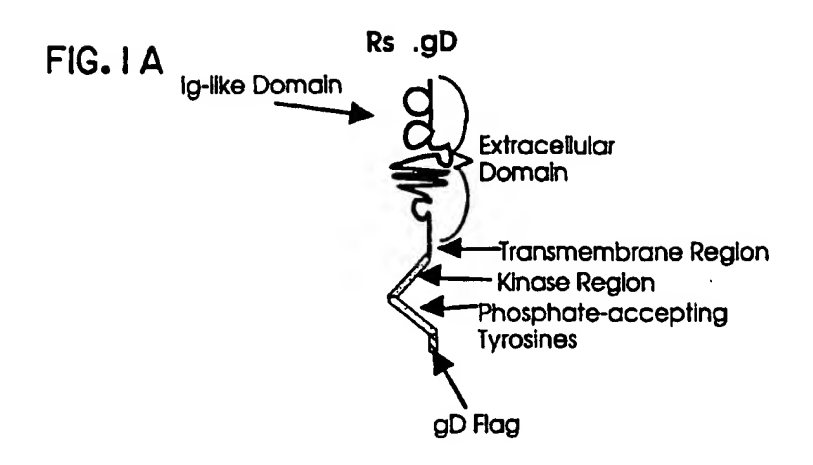
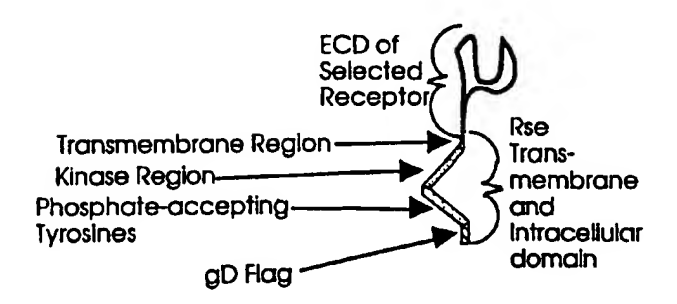
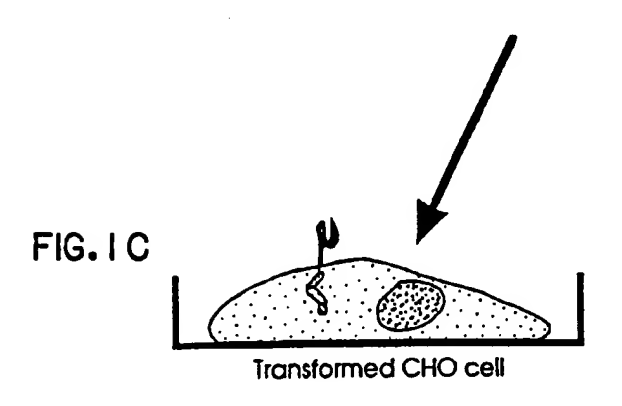


FIG. I B Receptor ECD/Rse.gD Chimera





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335 DSGLIEWEEVIPPEAP CCCCCTGGAGGCCCCTGGAAGCCCCTACAAACTG TCCTGGGTTC AAGACAATGG K G L A P A S A P Q N L H A I R T CAAGGTCTA GCCCCAGCCA GCGCTCCCA AAACCTCCAT GCCATCCGCA T V H L Q A L P A A P F N I T V T CACTGITCAC CITCAAGCAC IGCCIGCAGC CCCTTCAAC AICACGIGA 235 K L S S S N A S V A W M P G A D G R A L L Q S C T V Q V T Q A P G 701 CAAAGCTTTC CAGCAAC GCTAGTGTG CCTGGTGCTGAT GGCCGAGCTC TGCTACAGTC CTGTACAGTT CAGGTGACA AGGCCCAAA FIG.2A A T N Y S L R V R C GCCACCAACT ACAGCCTCAG GGTGCGCTGT 68 G M E E P D I Q W V K D G A V V Q N L D Q L Y I P V S E Q H W I G 201 gggatggag gagcctgaca tccagtgggt gaaggatggg gctgtggtcc agaacttgga ccagttgtac atcccagtca gcgagcagca ctggattgg GTGTGGCTCA GTCCCCCTGA S Q G Q P V K L N C S V E TGTCTCAGGG GCAGCCGGTG AAGCTCAACT GCAGTGTGA F L S L K S V E R S D A G R Y W C Q V E D G G E T E I S Q P TTCCTCAGCC TGAAGTCAGT GGAGCGCCTCT GACGCGCCC GGTACTGGTG CCAGGTGGAG GATGGGGGTG AAACCGAGAT CTCCCAGCCA E A V G GAGGCTGTGG GGCGGCTCTG P P N A P F Q L S C CCACCCAATG CCCCTTTCCA ACTGTCTTGT CCGCGCCTCG GGCTGCTGCT CCTGGTGCCT ď > \* ~ J TGCTCCGGGA CGCTGCCGCT GCCGCCGCCA K L T V AAGCTGACAG Ω GGGGGACCCG CCTTTCAGAC F T C L R T A CTCGCACAGC L A V TCTGGCAGTG AGCCCCGGTG ₽ 0 ວວວວເ F S C E A H N L K G L A S S TITTCCTGTG AAGCTCACAA CCTAAAAGGC CTGGCCTCTT 135 V E G V P F F T V E P K D 401 CGGTAGAAGG TGTGCCATTT TTCACAGTGG AGCCAAAAGA 168 P V T I V W W R G T T K I 501 ACCTGTTACC ATTGTCTGGT GGAGGGAAC TACGAAGATC GCAGGTCTGA AGCTCATGGG extracellular domain > 268 G W E V L A V V V P V P B01 AGGCTGGAA GTCCTGGCTG TTGTGGTCC TGTGCC 3 Ω 노 35 L P E S A A 101 TGCTCCGGA GTCCGCCGC \* « \* « G seduence L signal 201 601 101 301

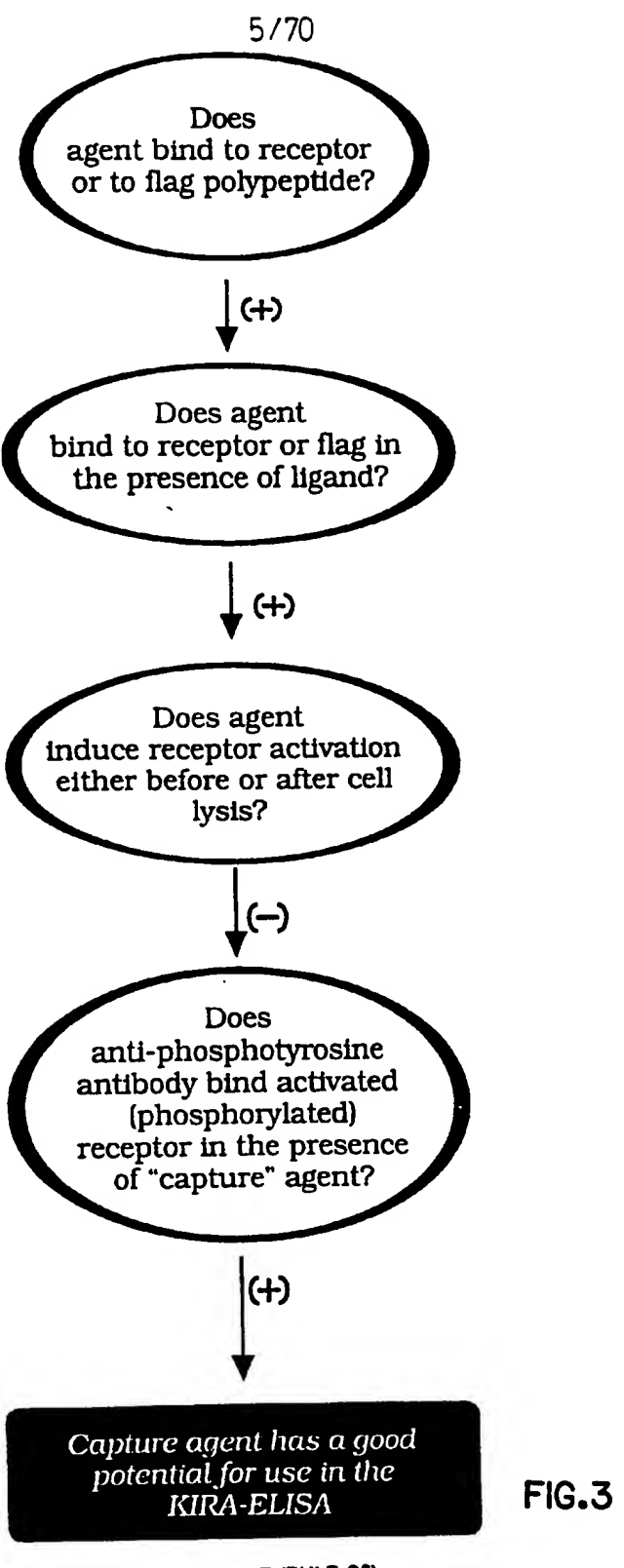
S V K L TCAGTGCGGG T L I R ACCCTGATCC CATCCCCATG CAGAGGACAT AGTTCCTCAG GTGGCTGGCC TTGACAGTGT CTTGGGGCATC ACAGCCGCAC ATCCTGGGTA CCTGTGGTCC CAATGCAGTT FIG 3 R A K G R L P AGGGCTAAAG GCCGTCTCCC GACATTGAAG C M L A TGCATGCTGG TGCCTGTCAA CATTGGACAG E F G AGAGTTTGGT F N L P L Q CCTTTAACCT ACCCCTCCAG G Q A F GGGCAAGCCT TGTGCGTCTC > ø H ф G K G TGGCCAAAGG Y S G D Y Y R Q G C A S K L TACAGTGGGG ACTACTATCG TCAAGGCTGT GCCTCCAAAC M L K A D I I A S S ATGCTGAAAG CTGACATCAT TGCCTCAAGC A R N TGCTCGGAAT R S F N R E R P E R I E A T CGGTCCTTCA ATCGAGAAAG GCCCGAGCGC ATCGAGGCCA D P Q K D L I V R V GGGATCCCCA AAAGGACCTG ATCGTACGTG CTTCGAAAGA GACGGAAAGA GACGCGGTTT œ intracellular domain D L A GAGACCTGGC A K L V G V S L R S GCCAAACTTG TTGGGTAAG CCTCCGGAGC G R M L G E N P G P P H S R I CTCCCGGATT TTCATCCACC GTTCACCCTG G Q Q AGGCCAGCAG Ö L L A TCCTGCTCGC CTCTCGGAAC A V K GGCTGTGAAG N L T G W CCAAT TTGACAGGCT E Q Q CAGAGCAGCA CCTCATCCTG ATGACCGTGC Ŋ 668 T V C V A D F G L S R K I 2001 GACAGTGTG TGGCTGACT TCGGACTCTC CCGGAAGATC 568 E A A C M K E F D H P H V 1701 GGAAGCAGCT TGCATGAAGG AGTTTGACCA TCCACACGTG V I L P F M K H G D L H A F GTCATCTTGC CCTTCATGAA GCATGGGGAC CTGCATGCCT 635 F M V D I A C G M E Y L S 1901 GGTTCATGGT GGACATTGCC TGCGGCATGG AGTACCTGAG 468 M A R G E P A V H F R A A 1401 CATGCCCCG GGAGAGCCAG CCGTTCACTT CCGGGCAGCC 535 A Q L K Q E D G S F V K V 1601 AGGCCCAGCT GAAGCAAGAG GATGGCTCCT TTGTGAAAGT 501 S D E L K E K L E D V L I P 1501 AGCGATGAAC TAAAGGAAAA ACTGGAGGAT GTGCTCATCC G C G P W S Q P L V V S S H GGCTGTGGAC CCTGGAGTCA GCCACTGGTG GTCTCTTCTC 368 T Q D E L T V E G T R A 1101 AACCCAGGAT GAGCTGACAG TGGAGGGGAC CAGGG 601 V 1801 GT 401 1201

FIG.2C 835 S G M G A V G G T P S D C R Y I L T P G G L A E Q P G Q A E H Q P 2501 gcagtggcat ggaggcagtg ggtggcactc ccagtgactg tcggtacata ctcacccc gagggctggc tgagccag gggcaggaggaggaggaggagg gD flag polypeptide
A D A S L K M A D P
GCAGATGCTA GCCTCAAGAT GGCTGATCCA 868 E S P L N E T Q R L L L L Q Q G L L P H S S C 2601 AGAGAGTCCC CTCAATGAGA CACAGGGCT TTTGCTGCTG CAGCAAGGGC TACTGCCACA CTCGAGCTGCT 901 N R F R G K D L P V L O 2701 AATCGATTCC GCGCAAAGA TCTTCCGGTC CTGTAGAAGC

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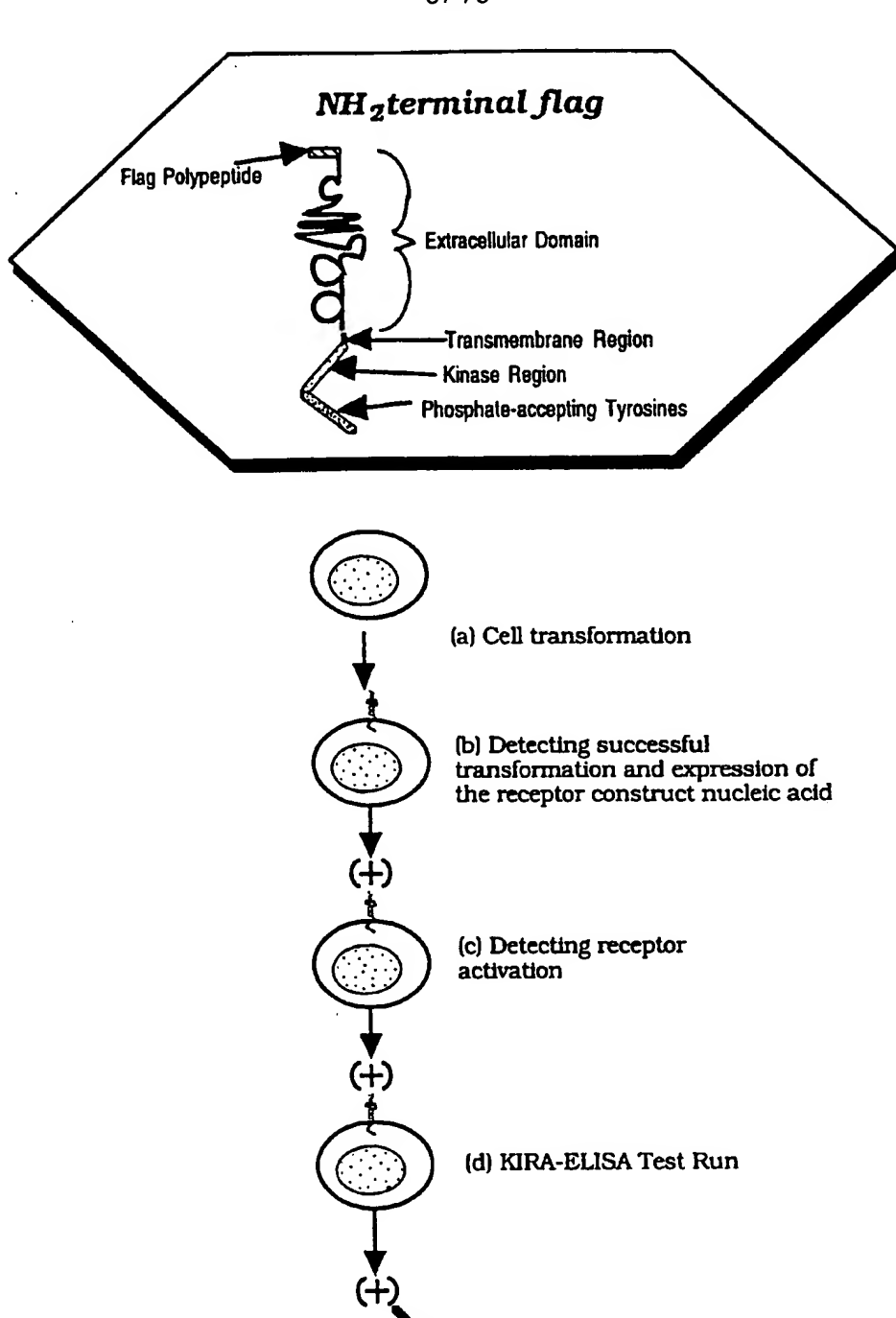
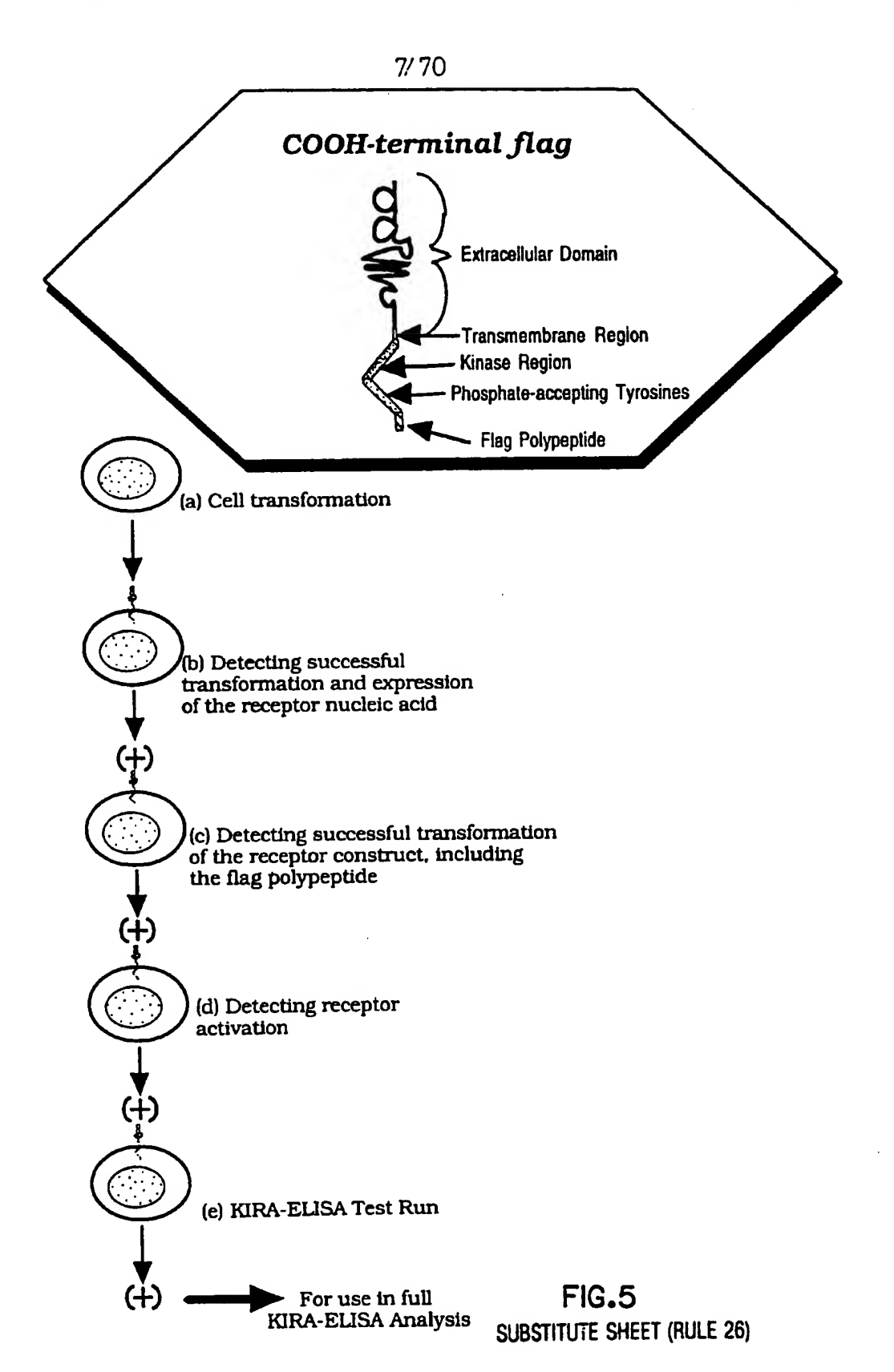


FIG.4
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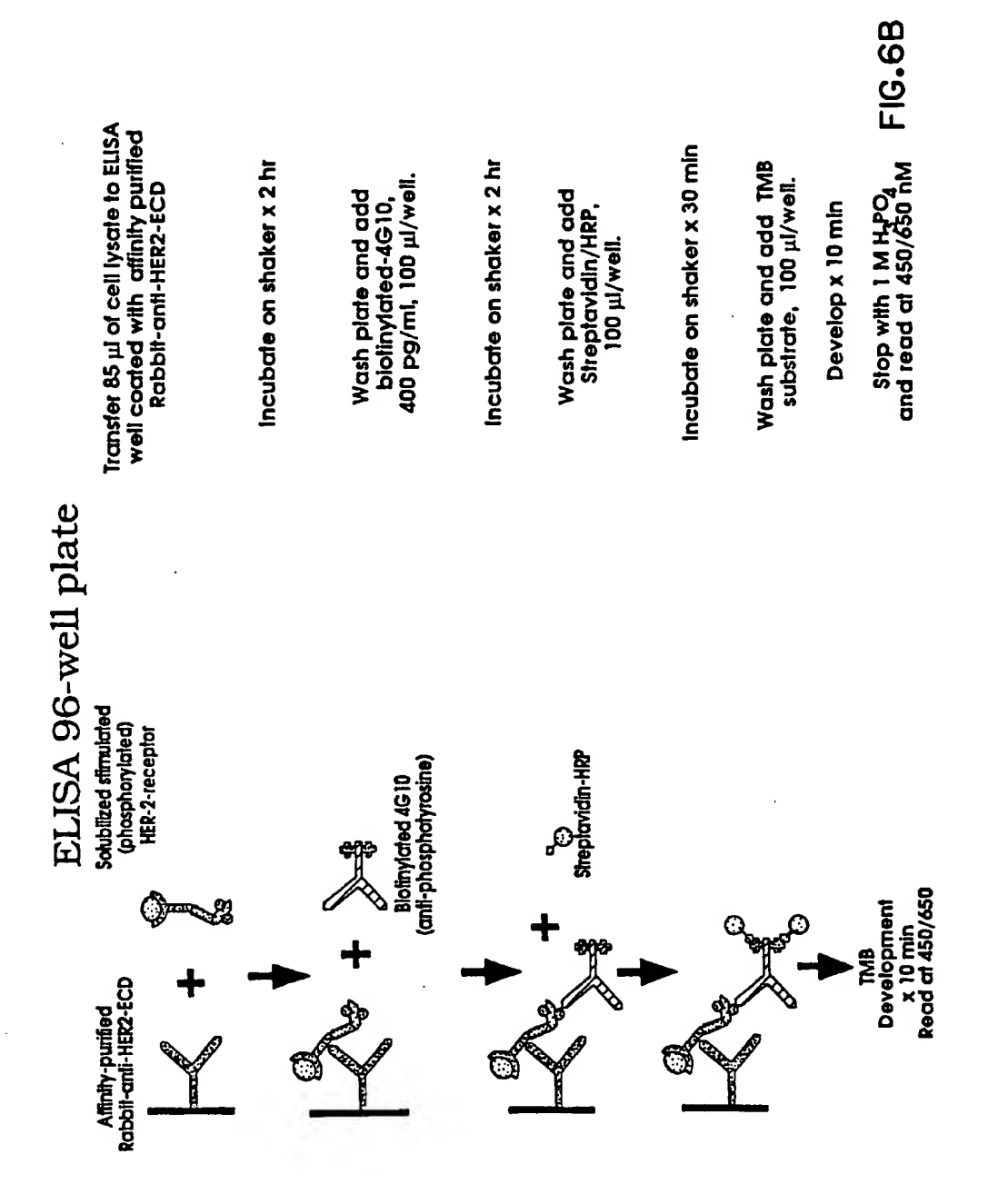
For use in full KIRA-ELISA Analysis

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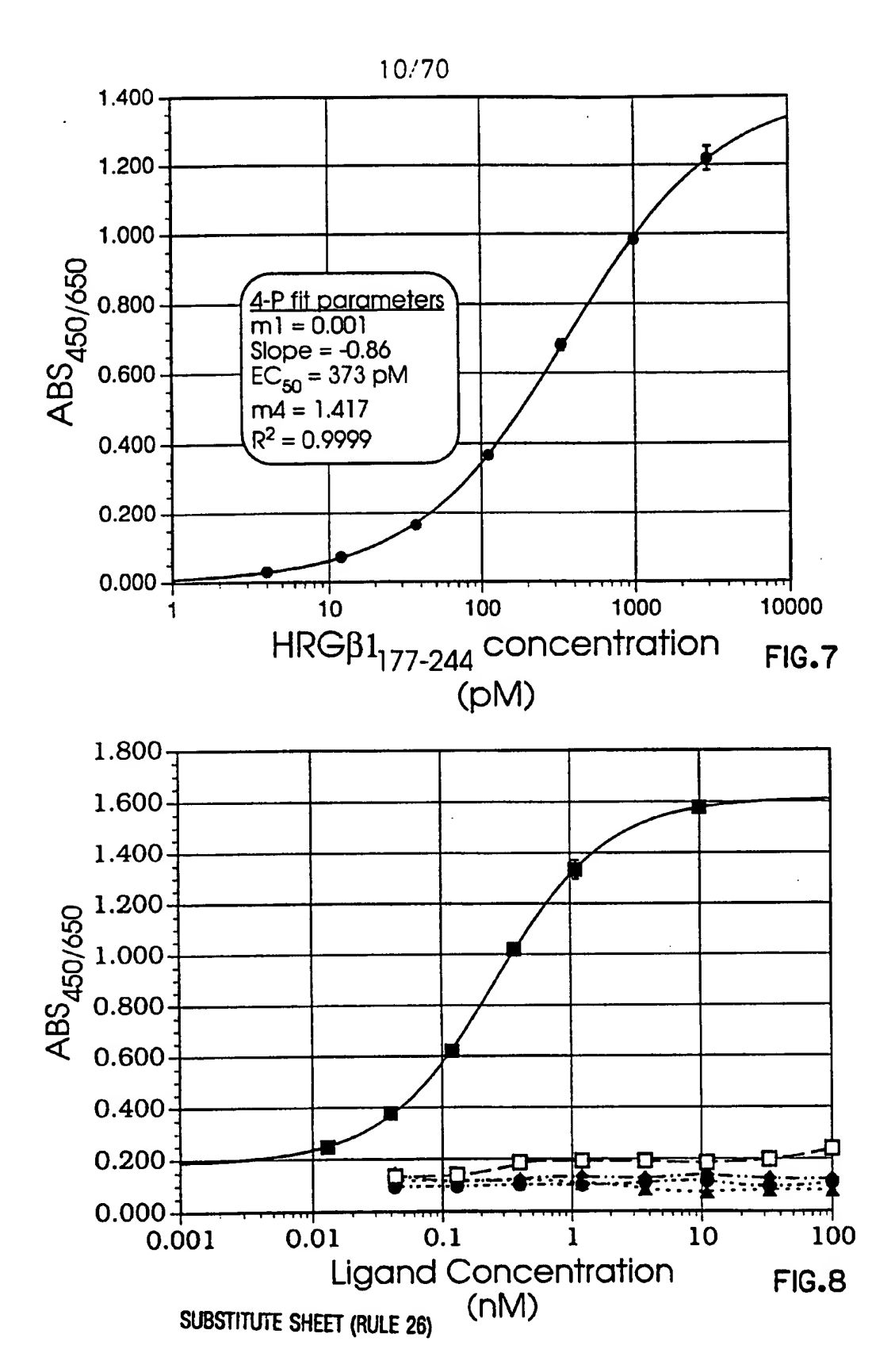
8/70 FIG.6A Na<sub>3</sub>VO<sub>4</sub> AEBSF, Leupeptin, & Aprotinin 2x10<sup>5</sup> MCF-7 cells per well in 96-well culture plates (flat-bottom) x 30 minutes at 37°C Stimulate with ligand. 50 µl per well 150 mM NaCi 0.5 % Triton-X 100 50 mM HEPES, pH 7.5 100 µl/well x 60 minutes Solubilize in: - Transmembrane Region HER-2 Receptor

Extracellular Domain Phosphate-accepting Tyrosines - Kinase Region **Phosphotyrosines** Heregulin SUBSTITUTE SHEET (RULE 26)

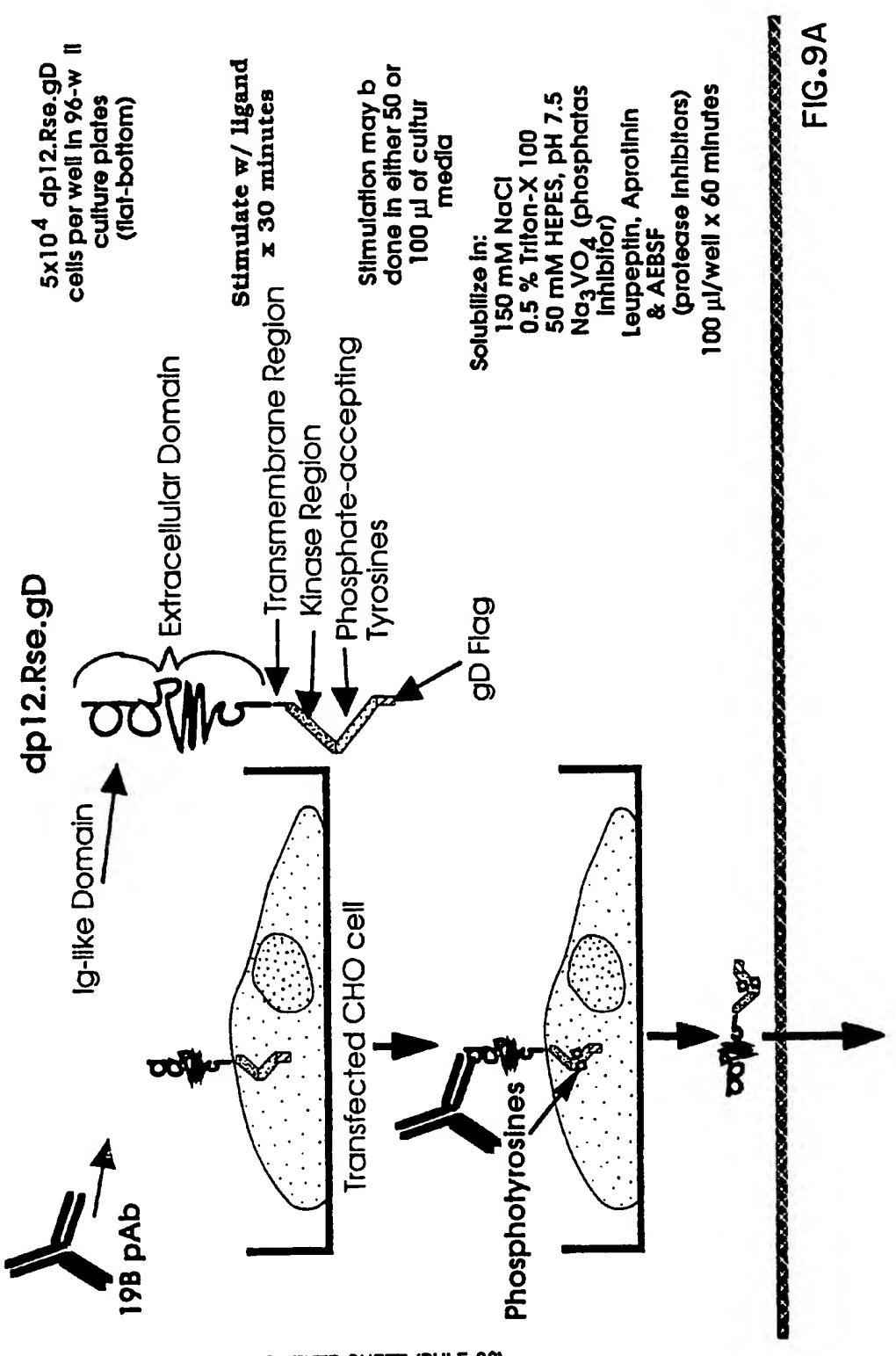


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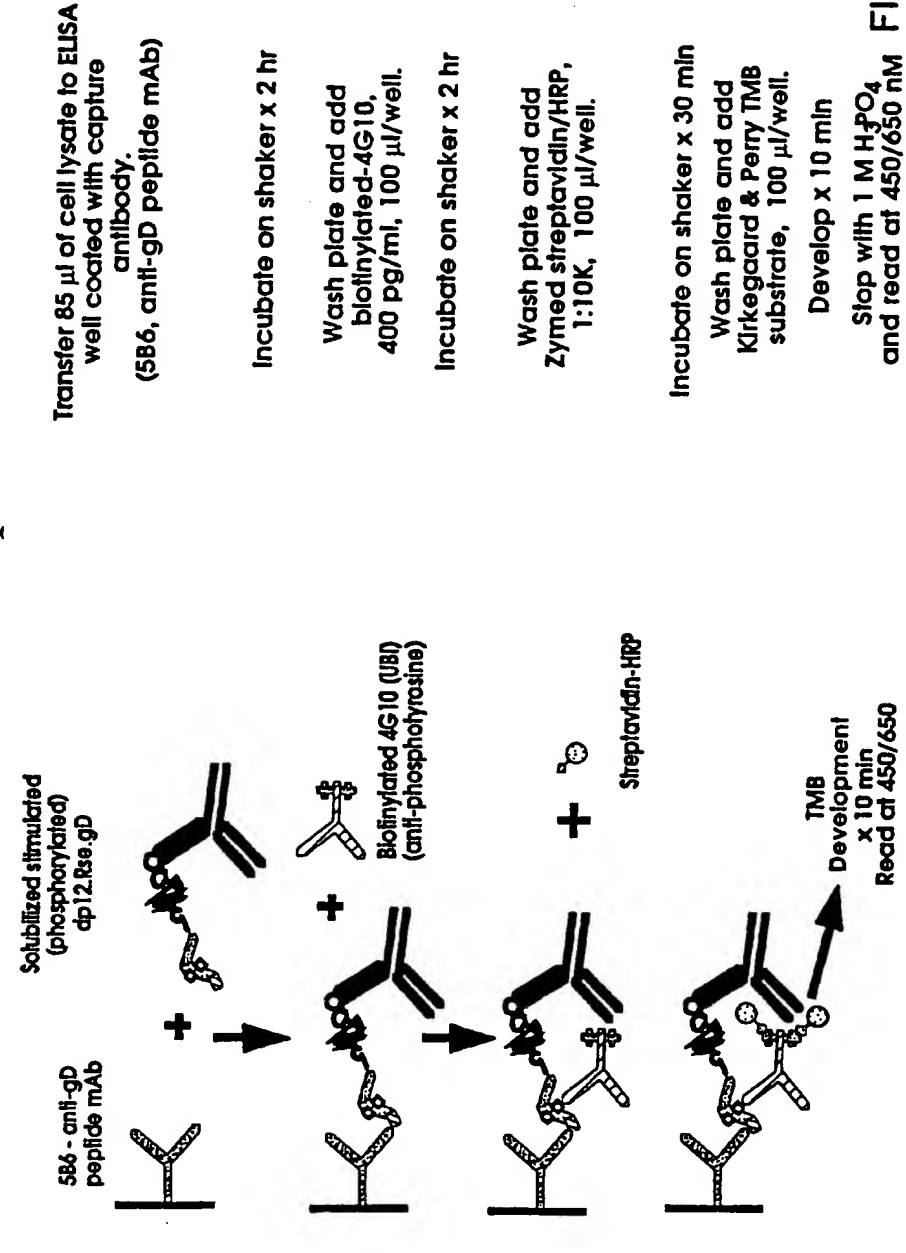
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FIG.9B

## ELISA 96-well plate



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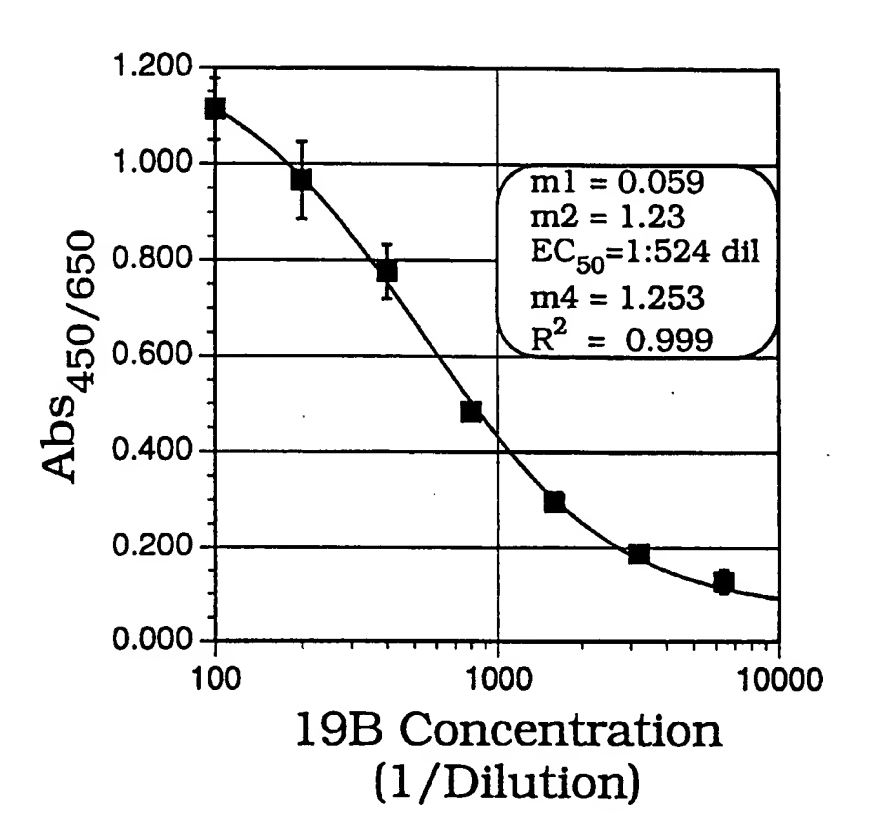
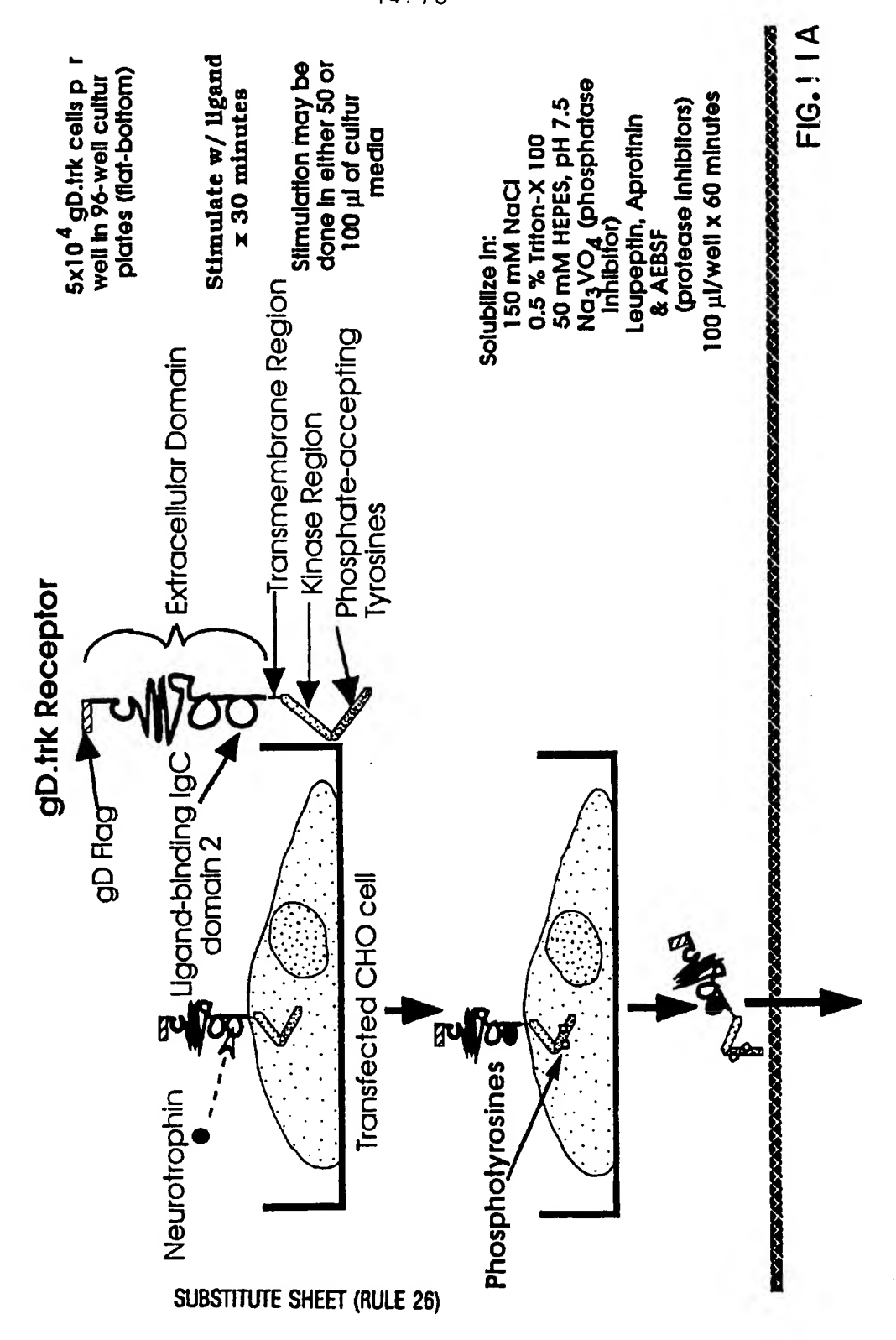


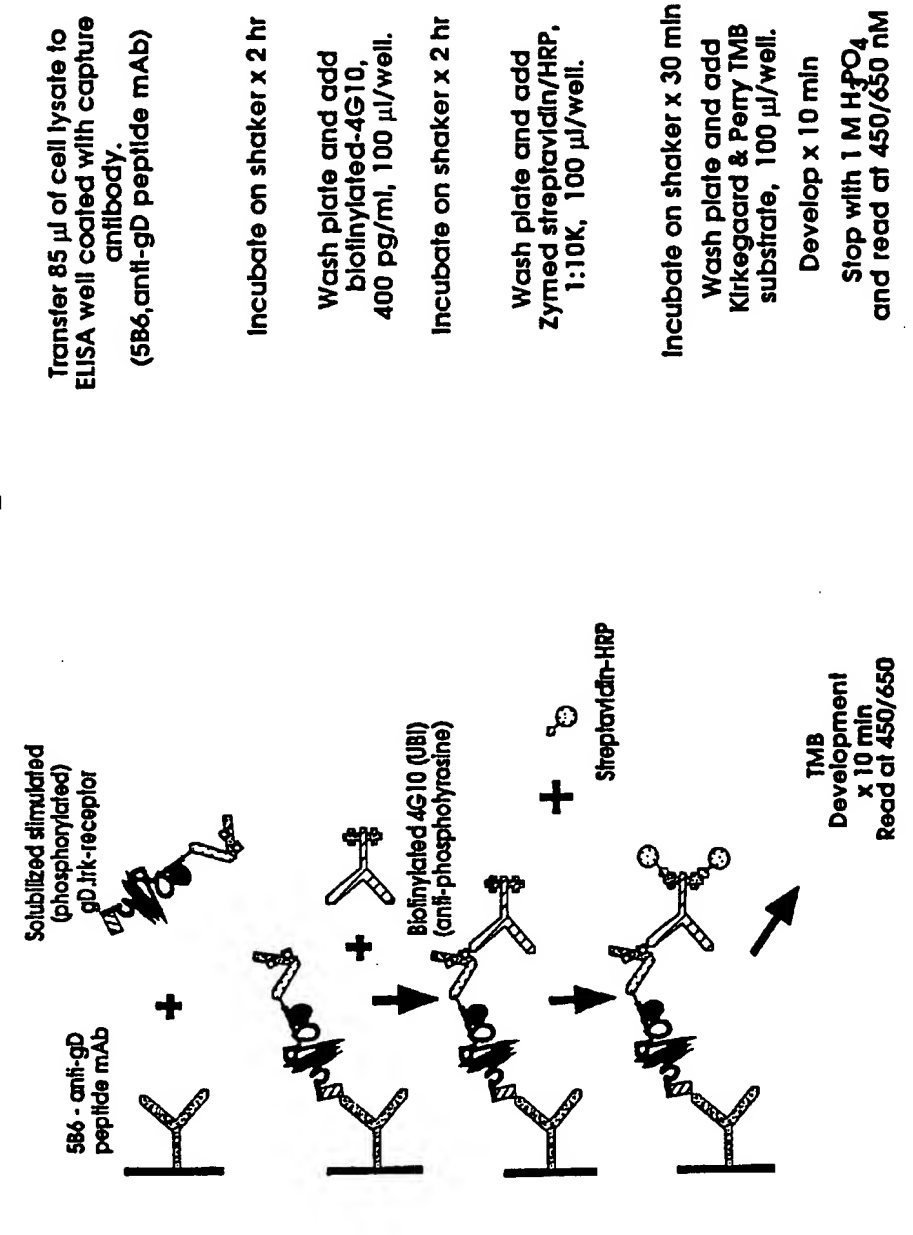
FIG. I O



<u>m</u>

FIG. 1

## ELISA 96-well plate



CAGGTTGACG GTCCAACTGC GGTGAGGGTC CCACTCCCAG CCACAGGTGT GGTGTCCACA CGGAAAGAGA GCCTTTCTCT GTAGGTGAAA TATAGAATAA CATCCACTTT start RNA ATATCTTATT 9ďs√ 841

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gD from pchadII AR1 site mutated in linker ^cloning

CCAGACATAG GGTCTGTATC CAGAGTCAGG GTCTCAGTCC ACGTCCTAGG TGCAGGATCC CACCAAGCTC GTGGTTCGAG GTGACGGAAG CACTGCCTTC TGGACTTAAG ^begin ACCTGAATTC 901

CGGAGAGTGT GCCTCTCACA TCCGAATATT ATTTGGGACG TAAACCCTGC TCAGGAACAG AGTCCTTGTC TGGCTCCAGT ACCGAGGTCA TTCCTGCTGG AAGGACGACC 961 SUBSTITUTE SHEET (RULE 26)

TTGCTGGTTG AACGACCAAC GAAGTCGCGC CTTCAGCGCG GACACTGTTC CTGTGACAAG TGACCCCTGG ACTGGGGACC CTCCGCGAGG GAGGCGCTCC AGAGCAGTTA TCTCGTCAAT 1021

g ^Start

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GTATGGGGGG CATACCCCC \* U GTGCGTTCCG CACGCAAGGC CTTTTGTGTG GAAAACACAC GAATTCCAGA CTTAAGGTCT TAGTCAATAG ATCAGTTATC ATGGGGCTAG TACCCCGATC 1081 H

\*> TCCATGGGGT AGGTACCCCA \*D \*H **1** TATCACCCGG ATAGTGGCC \*D \*A \*I CAAACAGCAG GTTTGTCGTC F\* V\* V\* CCGTGATTTT V\* I\* L\* GGCACTAAAA R\* L\* G\* A\* AGGTTGGGGG TCCAACCCCC CTGACGGCGG GACTGCCGCC T\* A\* A\* 1141 4

区 Ŀ M Z Ц Ω A Σ × 1 ß 4 d Y\* A\* L\* A R\* G\* K\* 24 1201

FIG. 12A ATCGATTTCG TAGCTAAAGC CGGCTGGGGT GCCGACCCCA TCTCAAGATG AGAGTTCTAC CGGATGCCTC GCCTACGGAG TATGCCTTGG ATACGGAACC GGCGCCGTTT CCGCGCCAAA

17/70

'Xho and GTA mutated in 'begin mature trkA

CGGGGCTACG GCCCCGATGC Ω GCCGCACCCT CGCCGTGGGA A CGAGCTCCAT GCTCGAGGTA > 回 L CTGGTCGA TGGACCAGCT O AC CTTCCGGTCC GAAGGCCAGG > GCCGTTTCTG CGGCAAAGAC P 1261

ACCTATCGGA TGGATAGCCT ഗ Д GATGGGGCCC CTACCCCGGG K G Д TACGTGGGCC ATGCACCCGG H U CGGGACTGCG CCTGACGC 只 H Ů S CACGGCTCCT GTGCCGAGGA S Ü 王 GACGACGGGG CTGCTGCCCC ρι C 64 1321

AGCAGCATCT TCGTCGTAGA O Ø ATCGAGAACC TAGCTCTTGG Z 回 H ACTCGAGATG TGAGCTCTAC × П 回 AGAACCTGAC TCTTGGACTG H CCCGGCGCAG GGGCCGCGTC 团 ъ G CCACCACCTG GGTGGTGGAC 니 н 1381 84

TCACCATCGT AGTGGTAGCA CTGAGAAACC H GACTCTTTGG Z L CCTGGGGGAG GGACCCCCTC 回 Ö Ы ATCTGAGGGG TAGACTCCCC Ü ĸ CTCGAGGCAC GAGCTCCGTG D r R 回 GCAGCATCTG CGTCGTAGAC H 二 Ø 104 1441

GGCTCAGTCG CCGAGTCAGC Н K TTCACTCCTC AAGTGAGGAG p, E Œ TGCCTTCCAT ACGGAAGGTA 工 [II K TGGCGCCAGA ACCGCGGTCT ሷ K CTCCGTTTCG GAGGCAAAGC > Ŀ 民 L CTTCTCACCA GAAGAGTGGT O ß 1501 124

TCCCGGAGAG AGGCCTCTC H Ö TTTTGACACG AAAACTGTGC Ø > H × AGAGAGGACC TCTCTCGG 3 Ŋ J CTCTGGAGTC GAGACCTCAG Ŋ 闰 TCCTTCAACG AGGAAGTTGC Z Ŀ ß GGACTTAGAG CCTGAATCTC H Z 144 1561

FIG. TGCGCTGGCT ACGCGACCGA 3 民 S C A L TCTTGTGCCC AGAACACGGG TCTGCACTGT AGACGTGACA U 二 H CGGGGAACCC GCCCCTTGGG Z CTGGTCCTGT ល GACCAGGACA H > Ы GAATGTCCTT CTTACAGGAA 闰 Ø ᆸ 164 1621

FIG.

ACGTGGTGAC

CGCCACCTCT

CGACGTGTGC

GGTCACACGT

AGGAAGGCC

CCAGTTGCAG

GGTCAACGTC TCCTTCCCGG

TGCACCACTG AGAGACAAGT TCTCTGTTCA ACTTCCAGGT AGGTGGAGGG TCCACCTCCC CCACGGTGAT GGTGCCACTA GTGACCTCAA CACTGGAGTT TGAAGGTCCA AGTGTCATGG TCACAGTACC 3 Ø Ø J H > > > 闰 H Ω > I × 터 S N V T S AATGTCACCA GACGCCACGG GAGCAGTCAG CGGCCAGAGG GCCCGTCTCC Σ GCGGTGGAGA CTCGTCAGTC L R C Q CTGCGGTGCC 4 TTACAGTGGT > CACGGGTGCG Ø GTCTTCGACG H GTGCCCACGC Q K L Q CAGAAGCTGC 回 S 团 Ö E > K Ø ρι 回 ĸ A > CTGGGACCGG GCTGCACACG GCTACACCCG CACAGAGCTG GTGTCTCGAC GACCCTGGCC CGATGTGGGC CGACGTGCTG GCTGCACGAC TCACGGACTT CAGCTGTGGT GTCGACACCA AGTGCCTGAA H J U Н Ø U 回 J > 团 出 > U Д Ы Ω E H Ω Ŋ > CCAGTGTGCA CCCTGGGGCT GCTGGATCCT GGGCAGAGAA CCCGTCTCTT TACACCCCCT CGACCTAGGA GGGACCCCGA ACGGGTTACG TGCCCAATGC ATGTGGGGGA CTGACCCGCC GACTGGGCGG Ø Н Ы Q > Н U 团 Z U H 3 K S E Q A G GAGCAGGCCG A S V D GCCTCGGTGG GGTCTGCCAT TTGACGTGCT CCAGACGGTA 3 AACTGCACGA L A H M CTGGCCCACA CTCGTCCGGC ß GACCGGGTGT CGGAGCCACC Σ GAGGAGGAGG CTCCTCCTCC C ပ Д 工 回 ᄓ H ĺΉ 回 ᆸ W හ GTCCTTCTTG CAGGAAGAAC CGTTCCCGGG CCACGGGTTA GAAATCTGGG CTTTAGACCC GGTGCCCAAT GCGGGGCCTG CGCCCGGAC GCAAGGGCCC TGTCGCGACC ACAGCGCTGG A N V U n u N d A Ωı ល ี . ผ DC, æ O 284 1981 304 2041 264 1921 1861 224 1801 244 1741 184 1681 204 SUBSTITUTE SHEET (RULE

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TCTTCAATGG AGAAGTTACC GACGCGACCG H CTGCGCTGGC 3 K CCGTGGCAGA GGCACCGTCT S Д TACCCGTCGG ATGGGCAGCC O U AAGAGACACC TTCTCTGTGG > Ŋ ſτι CACGTAGGGG GTGCATCCCC М 324 2101

GTCGGTTACT CAGCCAATGA Z K CTGGAGCCGG GACCTCGGCC A ሷ 臼 H GTGACTCAAG CACTGAGTTC ĮΣų 回 H GCTTCATCTT CGAAGTAGAA H נצו S TTACTCTGGT AATGAGACCA 回 Z GAGGCACGAG CTCCGTGCTC H > 344

ACGCCAACTA TGCCGTTGAT Z Ö GTGCAGTTGT Z CACGTCAACA Z > 出 GGTCGGGTGG CCAGCCCACC ф  $\mathbf{O}$ TGCGCCTCAA ACGCGGAGTT Z Ц K Ц GTGCCCACAG CACGGGTGTC Ö G 出 CTGGCACGCC GACCGTGCGG > 364 2221

CTGCCTTCAT GACGGAAGTA S I M A TCCATCATGG AGGTAGTACC GGCCTCCGCC CCGGAGGCGG K K CCTTCGGCCA GGAAGCCGGT Q Ö [14 GCTGCCAACC Д CGACGCTTGG z Ø GTGCGACGAC CACGCTGCTG H T I 384 2281

GCACATCTGG CGTGTAGACC ß H GACACTAACA CTGTGATTGT Ŋ Z H Д CCCCATCCCT GGGGTAGGGA Д Н ACCCCGAGGA TGGGGCTCCT Д 臼 ρ Z AAGCTCAAGT TTCGAGTTCA 阳 ſτι GGACAACCCT CCTGTTGGGA Д Z Ω 404 2341

TGGGCCTGGC ACCCGGACCG O > TCGGTGGCTG AGCCACCGAC > Ŋ TTTTGGGGTC AAAACCCCAG ^begin TM > O **[24** ACGAAACACC TGCTTTGTGG H 回 CTCTTCTTCC П GAGAAGAAGG × 回 AGACCCGGTG TCTGGGCCAC > Д Ω 424 2401

FIG. GTGGACGGAG CACCTGCCTC U CTCAACAAAT ŢΥ GAGTTGTTTA \end × Z H GCTCCTTGTG CGAGGAACAC > H H TCTACGCT AAAGATGCGA H H Ŋ TT H ACGGAGAAGG TGCCTCTTCC H U GCAGAAACGG CGTCTTTGCC K H > 444 2461

2E

FIG.

GCCGCCCCT

TGCACCGAGG

CTTCGGCGTC

TCGTGCGCTT

CACCAGCACA

GTGGTCGTGT

GTACGACGTC

CATGCTGCAG

AGCACGCGAA

GAAGCCGCAG

ACGTGGCTCC

CGGCGGGGA

20.770

GACAGTTCCG AGCTGCTCAC TCGACGAGTG TTCCGAGACC TTCACCACAT GGAAGGTCTT CCTTCCAGAA CTGTCAAGGC ACCGGTACAG AAGGCTCTGG AAGTGGTGTA TGGCCATGTC H Н Д H 노 > 工 Σ W > H × 足 田 U ATGCTGGTGG Ü GGCGCCTTTG CCGCGGAAAC A TACGACCACC 回 CGTGAGGCTG Q GCACTCCGAC TGGCTCCCGT > GATGCCTGTG CTACGGACAC CTCCTACCCG X L ACCGAGGGCA GAGGATGGGC > ĺΉ 回 U Ø U U IJ H 团 æ 回 Ø Д Σ Ω Ö C 召  $\vdash$ 回 CGTCCTGTTC GGACTTCCAA ATACTTCAGT CGACCCCCTC GCAGGACAAG CCTGAAGGTT GCTGGGGGAG GGACAGGGG TATGAAGTCA GCTGGCTCCA CGACCGAGGT CCTGTCCCCC > 回 × Ø S ф Ω Ĺ U O Ŀı Ŋ Д H **>**4 Õ [L4 H H TCCTGCCTGA AGGACGGACT GTGCTCGGCA CACGAGCCGT AGTTCACCCT AGAACCCACA TCAAGTGGGA TCTTGGGTGT GTGGCAGCTC CACCGTCGAG GCCCGGCTGT CGGGCCGACA Ø H 团 Ø ß Д 3 ĸ Д ល 노 Ц > Ø Z Ç Дı CGCAGGCTCT TGCCACAACC GCGTCCGAGA П CTGTAGCACG CHNL ACGGTGTTGG Ŋ H GACATCGTGC ATGACATTGG 回 CACATCATCG U GTGTAGTAGC CCCTAGTTGG TACTGTAACC GGGATCAACC D I O 田 团 H Н Z Ø Ŋ H H H I 田 Ø U GTTCGCGGCC TGACTTCCTC CGAGGTTCCG GGAACGACTC CCTTGCTGAG ACTGAAGGAG GGACGTAAAG CAAGCGCCGG GCTCCAAGGC TTTGTTCAAA CCTGCATTTC AAACAAGTTT ø 回 回 Ü X 民 Ŀ Ŀ ы L A Ø × Ξ J Σ H 584 2881 544 2761 564 2821 524 2701 504 2641 484 2581 464 2521 SUBSTITUTE SHEET (RULE 26)

FIG. 12F GTGTGGGAGG CACACCCTCC GTAAGTTCAC CATTCAAGTG ACCACCAGTT CGGGTCTGCA GCCCAGACGT TGGTGGTCAA CCTACGGCAA GGATGCCGTT CTAGGGTACC TGGGTCTGGG ACCCAGACCC GATCCCATGG 回 U ſι > Ö J Ц 民 × > > × U O Ü CTCTAGAAGT D Y Y R GACTATTACC CTGATAATGG K ATCCTGTACC TAGGACATGG H GAGATCTTCA Ø CACATGGACC CCGGTCCCTG K GCGAAGGAGG H GGCCAGGGAC CCAGGCCCCC GGTCCGGGGG GTGTACCTGG H K ĸ CGCTTCCTCC 니 U >1 [14 Д **>** H J × Н h  $\alpha$ H U 闰 Н U  $\mathbf{O}$ > ρι GATGTCGTGG CGGGCTCTCG GGTGCTCTGG CTACAGCACC CCACGAGACC GCCCGAGAGC CCTGGAGTTG CCTACACCGA TGCGGGGATG ACCCCCTAC CTGTCTAGTG GACAGATCAC GGATGTGGCT GGACCTCAAC S A H 3 Σ > K h Ч 回 Ŋ > Ö H Д > U K Д × Д 4 GCAGGGATAT GCTTCGGCGT GCTGGATGCC CGACCTACGG CGAAGCCGCA GCCAGGTCGC CGGTCCAGCG CCACACGCAA GGTGTGCGTT CGTCCCTATA GACCACCCT ACGCCGTGCC CTGGTGGGGA TGCGGCACGG ሷ > Н 团 K Z 闰 Σ > Ø Д U H ĸ <u>ෆ</u> 工 <sub>ල</sub> 3 [24 E 区 Z K AAACCGTACT K GACGGGTAAG S CTGCACACCT Ŋ CGGGACCTGG CTGCCCATTC GACGTGTGGA GCCCTGGACC S A V A S GCCGTGGCTA TTTGGCATGA Z TTCGACGACC S CGGCACCGAT AAACTCATAT K AAGCTGCTGG TTTGAGTATA Σ H d J 3 H J × K A Ц > Ö Ø Д 团 H **14** Ŀ  $\succ$ × [t4 AAAACACGTG GATTGGTGAT GGCGTGGTAC TTTTGTGCAC CTAACCACTA CACCGAGAGC CGTCGACGAC CCGCACCATG GTGGCTCTCG Q L L GCAGCTGCTG CGAGTACCAG TGGACTACGG ACCTGATGCC GCTCATGGTC Д 工 ល 3 T > > Ö 回 A Д Σ H [z, ĸ Н Ö ρι 724 664 3121 3181 704 3301 744 644 3061 684 3241 624 3001 604 2941

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FIG. 12G CGTAGTGTTT CTACCGGCGG GCATCACAAA GCTGCTGGCA CGACGACCGT TTCGGGACCG GATGGCCGCC AGGGACGTGA TCCCTGCACT AAGCCCTGGC Ø J 3 U A ATTTCGTTAT ATCATGCGGG Ø CGGCCGACG ATTAAGTTAG TAAAGCAATA GCCGGCTGC ACGTAGTGCG Ŋ TAGTACGCCC TAATTCAATC TGCATCACGC J ĸ ĸ Σ A Н TGGTTACAAA ACCAATGTTT GGGCTAGAAT GGTCTACGCC CCAGATGCGG CCTACACGTG CCCGATCTTA CCGTTAGCTG GGATGTGCAC GGCAATCGAC and fill^ Ø 王 primer^ > × Ω > site added with cloning with cut CCAGGCACCT CCTGTCTACC TGGATGTCCT CGAATATT GGACAGATGG ACCTACAGGA CAGCTTATAA ACAGCATCAA TGTCGTAGTT GGTTGTGCCT GCCCACCAGA CGGGTGGTCT CCAACACGGA J 团 × > H  $\Omega_4$ Ω ഗ ы GT site removed early poly AACAAATAAC ATGGCCCAAC TTGTTTATTG GTCGTTGCGG CCACGTGCCT GGTGCACGGA 工 CAGCAACGCC TACCAGCTCT ATGGTCGAGA U × ĸ Ø 民 > Q ^sv40 գ д R1 GGTCCGTGGA CGCCCTCGGG TACCGGGTTG CAACCTCGCC GCGGGAGCCC CGTCGGGACC GCAGCCCTGG GTTGGAGCGG R1 Д р 回 回 L 吆 804 3541 3601 784 764 3421 3481 3361 SUBSTITUTE SHEET (RULE 26)

CAGGTTGACG

GGTGAGGGTC

CCACTCCCAG

CCACAGGTGT

GCCTTTCTCT

CGGAAAGAGA

GTAGGTGAAA

ATATCTTATT

TATAGAATAA

841

CATCCACTTT

start

^sp6 RNA

GGTGTCCACA

GTCCAACTGC

F16.1 CTACCGGCTG GATGGCCGAC A 4 X GGAGAGAGTT CCTCTCTCAA X ı, S TTGGCGGATG AACCGCCTAC P Ω 4 **L**\* CAAATATGCC GTTTATACGG A\* K\* Y\* GGGTCCGCGG CCCAGGCGCC \* U \* \*^ G\* L\* H\* G\* GGCCTCCATG CCGGAGGTAC

CGTCATAGTG GCAGTATCAC CAAGCTTCAG TGTGGTGCGT ACACCACGCA GACGAGGCTT GTTCGAAGTC CTGCTCCGAA ATCCCAGAGT TAGGGTCTCA \* H \*^ GTCTCTTTG TTTTGTTTGT AAAACAAACA CTGGGACACT L\* F\* V\* CAGAGAAAAC TGTCATTTGG GACCCTGTGA CGAGACGTCC ACAGTAAACC GCTCTGCAGG A\* V\* I\* CCCGGCACT GGGCCGTGA TATCCTTAAG CTCCTGACCC ATAGGAATTC GAGGACTGGG GTCAAGTCCT CAGTTCAGGA CTTCCACCAA GAAGGTGGTT from pchadII \* U GATCATCAGT CGCCAGGTTG GCGGTCCAAC TAGAGGCG ATTCCACTGC CTGGTGGCTC CAATCTCCGC CTAGTAGTCA GACCACCGAG TAAGGTGACG **\***1 A\* R\* gD ^begin GT CCCCCTGACG GTTGATGGGG ATAGCTAACT CAACTACCCC GGGGGACTGC CACATCTCGT GTGTAGAGCA TATCGATTGA TATCTTCCTG ATAGAAGGAC \* E \* U ^cloning linker Start gD M\* G\* AGGCCATACC GCGCTTGCTG TCCGGTATGG CGCGAACGAC TGGAGCCAAG CAGGGGTCTG GTCCCCAGAC TATTGCCTCT ATAACGGAGA ACCTCGGTTC 1141 1201 19 1021 1081 961 901

in^ GTA mutated and xho

trkB^ mature start O H

CCATACAGGG GGTATGTCCC TCGACGAGCT AGCTGCTCGA GTCCTGGACC CAGGACCTGG AGACCTTCCG TTCGCGGCAA O CCCAATCGAT 1261 39

TCTGGAAGGC AAGCGCCGTT GGGTTAGCTA

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TGGCATCGTG ACCGTAGCAC ACCCTTCTCC TGGGAAGAGG ρ, TGGTGCAGCG ACCACGTCGC S CTCTCGGATC GAGAGCCTAG  $\boldsymbol{\vdash}$ 只  $\Omega$ AATGCAGTGC TTACGTCACG ഗ U TGCAGGACGT × ACGTCCTGCA U Ŋ 59 1321

AATTTTCATC TTAAAAGTAG Н TGTAGTGGCT ACATCACCGA 田 H  $\mathbf{H}$ CTAGGACTCT Z GATCCTGAGA 回 Д Д ATTGTCACAT TAACAGTGTA > S Z GATTGGAGCC CTAACCTCGG ρι 回 H CGTAAAGGCT GCATTTCCGA ĸ ρı Ŀ A 1381 79

TGTGGGACTG U > TTGAAGCTTA > Ø 回 GAAGATGATG > Ω A 回 AATCATCAAC Z H H AAAGGTTAGA 回 吖 × A N Q K GCAAACCAGA O 99 1441

ACACCCTGAC

AACTTCGAAT

CTTCTACTAC

TTAGTAGTTG

TTTCCAATCT

CGTTTGGTCT

TAAAGACTTT ATTTCTGAAA H ſΞι CTCATAAAGC GAGTATTTCG 4 × 二 AAATTTGTGG A TTTAAACACC > Ŀ TTCTGGATTA GACCTAAT h O AA S CAATTGTGGA GTTAACACCT A > Н TCTTTAGACT H AGAAATCTGA J R N 119 1501

CAGATCCTTT GTCTAGGAAA 召 S TGACGAGTTT ACTGCTCAAA S H CGAAACAAAC H GCTTTGTTTG × Z 民 GTTAAAATGG CAATTTTACC H ſιμ Z TGCAGCACAT ACGTCGTGTA H 耳 Ø AACAGCAACC TTGTCGTTGG H Z ഗ Z 139 1561

FIG. ATGTACGAGG TACATGCTCC O H GCAATCCATT CGTTAGGTAA ſτι ρι Z ATCCTGGTGG TAGGACCACC O > ы H CAGACTTGAC GTCTGAACTG H 回 Ŋ ACCTTGACTT TGGAACTGAA H Д Ы GTAAAGGCAG CATTTCCGTC 工 民 ſΞŧ 工 159 1621

CACTCAGGAT GTGAGTCCTA GGTCAGGTCT CCAGTCCAGA GAGGCTAAAT CTCCGATTTA 臼 CTGAGAGGTT GACTCTCCAA H ACACCTAGTT TGTGGATCAA ACACTGTAAT TGTGACATTA D 179

GATACCCAAT CTATGGGTTA D, Н GTTTGGACGT CAAACCTGCA Ø J Z TAAGGGGACC ATTCCCCTGG A 니 ρı CAGCAAGAAT GTCGTTCTTA Z ល TGAATGAAAG ACTTACTTTC 田 Z TTGTACTGCC AACATGACGG L U ≯ 1741 199

AGGAAAGTCT TCCTTTCAGA × Ü GACACCTCCT CTGTGGAGGA 团 团 > GGATTGGAGT H CCTAACCTCA J Z д AGACCGGCGT TCTGGCCGCA 4 K L GTAGACGTTT CATCTGCAAA Z K ß ACACCAAACG Д TGTGGTTTGC H U U 1801 219

GGATGTTGGT CCTACAACCA > Д ATATGTATTG TATACATAAC 3 × Σ Z CCGGTTCCTA GGCCAAGGAT д > Д GGCAGGTGAT CCGTCCACTA D U K CCTGTAGTGT GGACATCACA > Ŋ Ö I T L S ATCACATTAT TAGTGTAATA 239 SUBSTITUTE SHEET (RULE 26)

AAGGATAACT TTCCTATTGA Н 吖 AGGCTCCTT TCCCGAGGAA H ß Ü Ö TCGGTGTGTG AGCCACACAC 단 工 Ŋ GAATGAAACA CTTACTTGT H 闰 Z CCAAACATAT GGTTTGTATA Σ 二 × N L V S AACCTGGTTT TTGGACCAAA N 259 1921

AGAACATCCT TCTTGTAGGA > П TGGCGGAAAA ACCGCCTTTT Z 回 R ATCTCTTGTG > TAGAGAACAC ပ ທ TGGGAAGCAG ACCCTTCGTC Q × U CCGATGACAG GGCTACTGTC S Ω Ω N I S S AACATTTCAT TTGTAAAGTA 279 1981

FIG. | ATTTCTCGAA TAAAGAGCTT H [I4 CAACTATCAC GTTGATAGTG H Н H H F A P CATTTTGCAC GTAAAACGTG CCTCACTGTG GGAGTGACAC > Ħ Ц TAAGACAGTT ATTCTGTCAA Z > S GAAGATCAAG Ω CTTCTAGTTC O Д D 299 2041

CCCAAAACCA GGGTTTTGGT ρι ACTITCCGTT TGAAAGGCAA X GGTAAGTGAC CCATTCACTG CTGGTGCATT GACCACGTAA 3 GTCTGGTGGT CAGACCACCA AGAGGTTGGA TCTCCAACCT ρι 2101

CTGTACTAAA GACATGATTT U CCAAATACAT GGTTTATGTA  $\boldsymbol{\vdash}$ × × L N E S TTGAATGAGT AACTTACTCA 回 CGGGGCAATA GCCCCGTTAT H Ü GGTTCTATAA CCAAGATATT × [z X O GCGCTTCAGT CGCGAAGTCA J A 339

AGGGTGAGTG TCCCACTCAC Д TCGACCTATT AGCTGGATAA Z Д L Ø CCGACGGAGG GGCTGCCTCC H U Ö CCTCATGGTG GGAGTACCAC 王 þ 网 GGTTAGTGTG CCAATCACAC H HN TATGTACAAT ATACATGTTA H > 耳 359 2221

ACTCTTTGTC TGAGAAACAG Ø × 回 TACCCTTCCT ATGGGAAGGA Д ×  $\mathfrak{O}$ AAGAATGAGT TTCTTACTCA × 闰 Z × TCTAATAGCC AGATTATCGG ø Н H GGGACTACAC CCCTGATGTG × А ATGAACAATG Ü TACTTGTTAC Z Z 379

AAATTATCCT TTTAATAGGA × Z GTGCAAACCC CACGTTTGGG Щ Z K ATTGACGATG U TAACTGCTAC Ω Ω Н GACCGGACCT CTGGCCTGGA U Д 3 ACTTCATGGG TGAAGTACCC U Σ TAAAGACGAG ATTTCTGCTC 工 S 399 2341

CACGAACAGA GTGCTTGTCT Z E TCGGGGACAC AGCCCCTGTG A G A N D I GCGAATGACA CGCTTACTGT D TGGAACTGCA ACCTTGACGT Ø H U TACTTCTAAT ATGAAGATTÄ × Ω ഠ GATGTAATTT CTACATTAAA × Н > Ω 419 2401

TCTCTCGGTC AGAGAGCCAG Ø H GTCGGGAACA CAGCCCTTGT 耳 回  $\alpha$ GATAAAACCG Ü CTATTTGGC H × A AGACGTCACT TCTGCAGTGA E > Д AGGGAAGGTG TCCCTTCCAC ß Дı TCATTACTTT AGTAATGAAA H 回 Z 439 2461

FIG. GCTGTTTCTG CGACAAAGAC þ H TTTTGGTAAT AAAACCATTA Ξ > H GGATTTTGCC H CCTAAAACGG U j. O GTCTGTGGTG CAGACACCAC > > Ŋ TGGTGATTGC ACCACTAACG 4 н > > ATACGACACC TATGCTGTGG > K × 459 2521

F

TATCAGCAAT ATAGTCGTTA Н CAGCCTCCGT GTCGGAGGCA > Ø TACTTTCCGG ATGAAAGGCC U GTTCAAACCG CAAGTTTGGC × GTTCTGTGAG CAAGACACTC GAATTCAACC CTTAAGTTGG 2581 479

TCCATCTTCT AGGTAGAAGA S Д CCTCATTGTG GGAGTAACAC Z S ATCTCCAATG U TAGAGGTTAC TGAGGTAGTG ACTCCATCAC 耳 王 J CTGCCAGCCC GACGGTCGGG GATGATGACT CTACTACTGA Ŋ Д Ω Ω 499 2641

CATTGAAAAT GTAACTTTTA 闰 Н AGATCCCTGT TCTAGGGACA > Д Н × CCTTACTGGT GGAATGACCA E Σ U TGTCALTALL ACAGTAATAA Н Н > GCCCAGATGC CGGGTCTACG K РО AGCCTTCCAC TCGGAAGGTG C Ö 回 Ø 2701 519

AGTCGTGTAG TCAGCACATC 耳 Ø ACACATTTGT TGTGTAAACA > Ŀ H О CTCAAGCCAG GAGTTCGGTC ы × GTTGTCAGTC CAACAGTCAG Q Ŋ Z AACCGTAGTG TTGGCATCAC H Н U CCCCAGTACT GGGGTCATGA Ľ × O Щ 5392761

TTTTCACAAG AAAAGTGTTC > × GAGCCTTTGG CTCGGAAACC U Ŀ 4 GATCCGCTTC U CTAGGCGAAG 臼 O CTTTTCCCTC GAAAAGGGAG 回 吖 노 ACATTGTTCT TGTAACAAGA J > Н TTCGCTGTAT AAGCGACATA Z H 559 2821

TCACTTCTGG AGTGAAGACC H × > TCTTGGTGGC AGAACCACCG Ø > H CAGGACAAGA GTCCTGTTCT H 봈 Д Ø CTGTCCTGAG GACAGGACTC 回 ф U GCTATAACCT CGATATTGGA J Z × CTAGCTGAAT GATCGACTTA U 回 Þ ᆸ 579 2881

FIG. 13E CGAGGACTGG GCTCCTGACC H H L CACTCCGGCT GTGAGGCCGA 回 Ø 回 GACTTCCACC ĸ CTGAAGGTGG 耳 Ŀ D TGCACGCAAG GTGCGTTC × 叫 AC A CCAGTGACAA GGTCACTGTT Z Ω ហ CTGAAGGATG GACTTCCTAC K Д LK 599 2941

D P L GCTGGGGGAG CGCACCTCCC GCGTGGAGGG TATGGCGTCT ATACCGCAGA ပ > GCAGTTCAAG CGTCAAGTTC > ATGAGCACAT TACTCGTGTA Н 闰 TTGGAGGTCG 耳 AACCTCCAGC O<sup>1</sup> Ч 619 3001

CCGTGTGCCG GGCACACGGC Ö 耳 4 TCAAGGAGTC AGTTCCTCAG 民 H ſτι CTGGAGTTGT X GACCTCAACA Z H Ω GAAGCATGGG CTTCGTACCC O 耳 봈 TTGAGTACAT AACTCATGTA × 闰 ATCATGGTCT TAGTACCAGA ∨ ਸ Σ H 639 3061

GTCGCAGATG CAGCGTCTAC 0 Ŋ AACTGACGCA TTGACTGCGT Ø H H CCGCCCACGG 团 GGCGGGTGCC E ρι Д TGAGGGCAAC ACTCCCGTTG Z Ö 回 ACGACTACCG TGCTGATGGC CCTGATGCCG GGACTACGGC > 659 3121

GCACTTCGTG CGTGAAGCAC ſΞι A S Q TGGCGTCCCA ACCGCAGGGT ATGGTCTACC H TACCAGATGG > Σ ອວວວອວອວອ ၁၅၅၁၁၁၁၁၁ U K CCCAGCAGAT GGGTCGTCTA Н œ Ø CTGCATATAG GACGTATATC Ø Н Ή h 679 3181

GAAAATCGGG CTTTTAGCCC Н × ACTTGCTGGT TGAACGACCA > H H GTCGGGGAGA CAGCCCCTCT Z 回 U GAACTGCCTG CTTGACGGAC H U Z CACCGCGATT TGGCCACCAG ACCGGTGGTC A T R GTGGCGCTAA R U L 二 699 3241

ACCGGTGTGT TGGCCACACA 二 U ACAGGGTCGG TGTCCCAGCC 
 Co
 > 14 ACTGACTACT TGACTGATGA × × A CGTGTACAGC GCACATGTCG Ŋ × > TGTCCCGGGA ACAGGGCCCT Д S R CTGAAACCCT Σ GACTTTGGGA U ſτι Ω 719 3301 SUBSTITUTE SHEET (RULE 26)

GTGCTGCCTT CACGACGGAA H H ACAGGAAATT TGTCCTTTAA ſt4 X 民 AGCATCATGT × TCGTAGTACA × Н Ŋ CGGAGGTCTC GCCTCCAGAG 臼 Д Д TTCGCTGGAT AAGCGACCTA Σ 3 ĸ ATGCTGCCCA TACGACGGGT H Д ᄓ Σ 739 3361

FIG. 13F CAAACAGCCC Д O × TCACCTATGG U × E TGGGAGATTT Ħ H 回 3 TCGTGTTG J > S > GGAGCCTGGG O h Ŋ AGCGACGTCT 3 > A Ŋ 3421

FIG. TGTTTATTTC ACAAATAAAG TATAATGGTT ATATTACCAA CCAACTTGTT TATTGCAGCT ATAACGTCGA GGTTGAACAA GGCGGTACCG CCGCCATGGC AGAAGCTTGG TCTTCGAACC

3841

GAGCTGGACG GGCTCTTCGA TAGTTTCTGA K early poly GTCATAATTG sv40 AAGTGAGACT CGACGAGAGG

ATCAAAGACT CCGAGAAGCT CTCGACCTGC from subcloning^ Sal site Xho half CAGTATTAAC half TTCACTCTGA GCTGCTCTCC 3781

TCCGGTGGTT GACGGCGACC GTAGAAAATT AGAGGATGAA TCTCCTACTT TCTGCCCGAC GCATGAGGAG CGTACTCCTC

AGGCCACCAA

CTGCCGCTGG

CATCTTTAA

AGACGGGCTG

TCCTTCCCAA AGGAAGGGTT CCCAGACCGA GGGTCTGGCT CCGGGAAAAG GGCCCTTTTC L G O TCTAGGCTAG AGATCCGATC ACCTGGACAT TGGACCTGTA S P V Y TCTCCGGTCT AGAGGCCAGA 839 3661

H Д Ы >

CCGGTTCCGT AAGTCTTGAA GTATGGGAGG GTTCCCGTAG CCTTCTTGTA GGGGTGTACT

GGCCAAGGCA K X Þ TTCAGAACTT H Z Ø H CATACCCTCC 니 H 工 CAAGGGCATC HU × GGAAGAACAT H × CCCCACATGA 叱 Σ 3601 819

GCAGCGAGAG CGTCGCTCTC TGGGGTGCTG ACCCCACGAC GAGCTGATGC CTCGACTACG GGAGGTGTAT CCTCCACATA CGTGCCCCCA GCACGGGGGT GCTGGGGCGT CGACCCCGCA 3541

闰 K Ø Z U U H Σ J 团 × > 回 O Д U H ĸ ρı PL, 799

TCAGGACGTC GAGTCCCGGC CTCACATAGT ACTCCACTAT ACAGTTTGTT ACCATGGTCG

AGTCCTGCAG GTTTGTCGGG AGTGGATACC CTCAGGGCCG K U Ø GAGTGTATCA ACCCTCTAAA H H U 回 CCAGCACAAC TGAGGTGATA 团 CCTCGGACCC TGGTACCAGC TGTCAAACAA Z Z H Ø ×

TCGCTGCAGA 779

CAGGTTGACG GTCCAACTGC GGTGAGGGTC CCACTCCCAG GGTGTCCACA CCACAGGTGT CGGAAAGAGA GCCTTTCTCT GTAGGTGAAA CATCCACTTT start ^sp6 RNA ATATCTTATT TATAGAATAA 841

in 'RI site mutated ^cloning linker

CCAGACATAG GGTCTGTATC CAGAGTCAGG GTCTCAGTCC ACGTCCTAGG TGCAGGATCC CACCAAGCTC ACCTGAATTC CACTGCCTTC GTGACGGAAG from pchadII TGGACTTAAG ^gD 901

CGGAGAGTGT GCCTCTCACA TCCGAATATT AGGCTTATAA ATTTGGGACG TAAACCCTGC AGTCCTTGTC CAGGAACAG H TGGCTCCAGT ACCGAGGTCA TTCCTGCTGG AAGGACGACC 961

TTGCTGGTTG AACGACCAAC CTTCAGCGCG GAAGTCGCGC GACACTGTTC CTGTGACAAG ACTGGGGGACC TGACCCCTGG CTCCGCGAGG GAGGCGCTCC TCTCGTCAAT AGAGCAGTTA 1021 SUBSTITUTE SHEET (RULE 26) ^Start

CATACCCCC GTATGGGGGG \*5 **\*** \* **X** CACGCAAGGC GTGCGTTCCG CTTTTGTGTG GAAAACACAC CTTAAGGTCT GAATTCCAGA TAGTCAATAG ATCAGTTATC ATGGGGCTAG TACCCCGATC

1081

g

AGGTACCCCA H\* G\* Λ\* TCCATGGGGT I\* A\* G\* L\* ATAGTGGGCC TATCACCCGG CAAACAGCAG GTTTGTCGTC F\* V\* V\* CCGTGATTTT GGCACTAAAA 1\* L\* \*> R\* L\* G\* A\* TCCAACCCCC AGGTTGGGGG CTGACGGCGG GACTGCCGCC A\* A\* \* E 1141 ゼ

FIG. 14A ATCGATTTCG TAGCTAAAGC Ŀ K CGGCTGGGGT Z GCCGACCCCA ρ, Ω A TCTCAAGATG AGAGTTCTAC × × L GCCTACGGAG CGGATGCCTC S A Q TATGCCTTGG ATACGGAACC Y\* A\* L\* A CCGCGGCAAA GGCGCCGTTT R\* G\* K\* 1201 24

trkc mutated Ų ^begin mature Z GTA Д and Ų site > 田 ^Xho 긔 O A > 괵

ATTGTGTCTG TAACACAGAC ACGGGACGTT TGCCCTGCAA CGAGCTCCAT GCTCGAGGTA ACCAGCT TGGTCGA ACC TGG GAAGGCCAGG CTTCCGGTCC GCCGTTTCTG CGGCAAAGAC 이 1261 44

CCCTCCTGGA GGGAGGACCT h ᆸ TTGGAGAAGG ሷ AACCTCTTCC [14 H Z GGACGATGGG CCTGCTACCC U Д Д 5522522 225225225 ρι 召 CGG ø GAGATCAATT CTCTAGTTAA U Z H 臼 GTCGTTCTGA CAGCAAGACT 댐 S 1321 64

ACATCTCAAG TGTAGAGTTC Ŋ H Ω AACATCACGG TTGTAGTGCC H Н Z GCGGTTATAG CGCCAATATC Н Z Ø CGTTACCCTT GCAATGGGAA Z U Z S G N S TCAGGGAACA AGTCCCTTGT S Z AGGCCAGGAT TCCCGTCCTA Ω Ø U 1381 84

ACCCCGTGGA TGCGGCACCT A > Ø GTGTGCGAGT ·Z CACACGCTCA H H H GCGCAGTCTT CGCGTCAGAA Ø K TAGAGAACTG ATCTCTTGAC Σ, Z 团 S I H I TCCATACACA AGGTATGTGT Н GAATATCACT CTTATAGTGA E Н Z 1441 104

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TTCGGAGCAT AAGCCTCGTA S ĸ H TTGAGTCCTG AACTCAGGAC Ö Ŋ Z GACCATCAAG CTGGTAGTTC 봈 Н H TTCAAAAGCT AAGTTTTCGA H × ø L TACACCGGAC ATGTGGCCTG O ¥ CATGGAGCTC GTACCTCGAG H 回 Σ 124 1501

CAAGTAACCG GTTCATTGGC Z ល ATAAACCTGT TATTTGGACA S H Z Н TTTGCGTTAT AAACGCAATA × П AGAACCCCCA TCTTGGGGGT 工 ρι Z X CGGAAACGGT GCCTTTGCCA K A F AGTCGGGTCT TCAGCCCAGA K Д Ø 144 1561

FIG. 14B TGCAGTTGGA [1] H Ø L R E L CTTCGGGAAT GACGCTGAGT S 니 E AGCTCTTCCA Ø 14 Ы Ø CTCTCGTGGC 3 Ŋ H GCTCACCACA E H ч 1621 164

ACGTCAACCT GAAGCCCTTA CTGCGACTCA TCGAGAAGGT GAGAGCACCG CGAGTGGTGT

AGGAGCAGGG TCCTCGTCCC O 回 GTCGAGACCG Q L W Q CAGCTCTGGC GGCGACCTAC CCGCTGGATG X 3 只 GTGACAT CACTGTA H Ω CGA( GCT U AAGTTGACGT TTCAACTGCA S U Z [±4 Q N F GCAGAACTTT CGTCTTGAAA [Li 1681 184

GGGTCGAAGG CCCAGCTTCC h Ø GCTGATGGCT S CGACTACCGA <u>ෆ</u> Ω Ø CTGCATCAAT GACGTAGTTA Z Н C TCTTGGAGAT AGAACCTCTA × ļ Z CTCAACAGCC GAGTTGTCGG Ø ഗ L N GGAGGCCAAG CCTCCGGTTC 봈 1741 204

CGGTGCAGTT GCCACGTCAA Z > 二 TAGTCGCACT S ATCAGCGTGA > Ŋ CCTTCCTGAG GGAAGGACTC 闰 Д GTCAGTGTGA CAGTCACACT Ö Ø ATGAACATCA TACTTGTAGT Ŋ Н M AGAGAAGGCG TCTCTTCCGC ĸ 년 기 1801 224

GATCACCCCT CTAGTGGGGA ρι Ŋ AATGGCTCTG TTACCGAGAC Ü S Ö Z ATAGTGAACG TATCACTTGC U H Н ACAATGCTGT TGTTACGACA > Z CGAGAGGGTG GCTCTCCCAC O D 风园 GGACTGGCAT CCTGACCGTA LTV SUBSTITUTE SHEET (RULE

AGACCAATCT TCTGGTTAGA H AACACTCACC Ö TTGTGAGTGG H ٤ Z GCAGTCCATC CGTCAGGTAG Н ß Ø TCACTGGGCT AGTGACCCGA H U > GACTGGATAG CTGACCTATC H M Q TCCTGATGTG AGGACTACAC P D V

GTGAGGACAA CACTCCTGTT Ω 回 TTACACTGCT S AATGTGACGA T T Z CTGCGACCAC GACGCTGGTG > u H CCATCAACTT GGTAGTTGAA u Z AATGTTCATG TTACAAGTAC 4 王 > N CTTGACCTGG GAACTGGACC E 3 284 1981

FIG. CCAGTGTTGC GGTCACAACG > S TACTCGTTAC ATGAGCAATG 4 Z S Σ GCACCACCCG CGTGGTGGGC Ö > > AACGTCTCTT TTGCAGAGAA Z 回 K L T C I CTGACGTGCA GACTGCACGT ACCGAAGTGG TGGCTTCACC 단 Ö 2041 304

TGCGCCTGGA

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ACGCGGACCT

CTCGGACTCG

CTCGGACCTC

GTGCACACCA

ATGATAGGGG

GGAGTGACAG

CCTCACTGTC

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TACTATCCCC

CACGTGTGGT

GAGCCTGGAG

GAGCCTGAGC

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GCCTGCACAA

ACGCTGCACT

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GGGGGGTGGT

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CTCAAACACC

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FIG. 14D CAGTTGGACT H U > GTATCCATAG H Ø TW > ^begin CACTTTTGGG ₽ CAGAAGAAGA A M 回 TATCACTGTG ACCCACAAAC Д I H > H 2461

ACTATACCCT TCCTCAAGGA AGGAGTTCCT CCACACCTCC GGTGTGGAGG AGGGAGAGAT TCCCTCTCTA TGATATGGGA E Ч AATGGCCACT TTGTTACCGT AACAATGGCA GAAGTGAGTC CTTCACTCAG TACTACCAAG ATGATGGTTC Z ſΉ TTACCGGTGA ecd insert Ħ O Ŋ U > Z Z 臼 GGTCTGGTAG GTGGGTGATG GAACAAACTG CCAGACCATC CCATGTGGAA GGTACACCTT CACCCACTAC CTTGTTTGAC ^end Н × E H ſι Ø ATAACTTTAT CCAAGATCAT TCAACAAGCC AGTTGTTCGG GCACAGCCAA CGTGTCGGTT TATTGAAATA GGTTCTAGTA Z ρ, H insert Ø Z H Z begin ecd GAGAGCACGG TGCCTGCTCT L R E S CTGCGGGAGT GACGCCCTCA TTGGGTGACC ACGGACGAGA U AACCCACTGG CTCTCGTGCC Ŀ 니 C. L. L 단 о, Z 团 TGGGCAGCCT AAGGCTCCCG GTAACGGTTT CGGGAAAGGT ACCCGTCGGA TTCCGAGGGC CATTGCCAAA GCCCTTTCCA × U 回 K Н 2401 404 424 444 384 2341 364 2281 SUBSTITUTE SHEET

GTCAACCTGA CATAGGTATC GTGAAAACCC GTCTTCTTCT TGGGTGTTTG ATAGTGACAC

AATATGGTCG TTATACCAGC × TW ^end ATGATCAACA TACTAGTTGT Z H Z AGAGAAGCAG TCTCTTCGTC > H H TGTTGGTGGT ACAACCACCA > Þ H H CGGACACAGG GCCTGTGTCC > ACGACGAAAA TGCTGCTTTT 2521 464

Ŋ Ω 曰 团 U ល Н > K > <u>p</u> <sub>ෆ</sub> × Σ O [I4 Ŋ

TCCTGAGTCG AGGACTCAGC AGTGGTGAGG GGCTGTCATC AGGGTCCCGT TTTGGAATGA ACGGTCCAAA 484 2581

TCACCACTCC CCGACAGTAG TCCCAGGGCA TGCCAGGTTT AAACCTTACT

TACGGCCCGG ATGCCGGGCC Ö Ø Д TCGTCACTGG AGCAGTGACC Ч S S CACCACGCCC GTGGTGCGGG Ω, ۲ E ACCACGGCAT TGGTGCCGTA  $\boldsymbol{\mathsf{H}}$ U CACCACATCA GTGGTGTAGT Z H 田 出 CAGCCCACTG GTCGGGTGAC H Д Ŋ 504 2641

AGTACTTCCG TCATGAAGGC × GAGAACCCCC CTCTTGGGGG Q щ Z 闰 CCCTGTCATT GGGACAGTAA H > D1 TGACTCGCAT ACTGAGCGTA Н CAGTAACCGT GTCATTGGCA E C T A GCTGTGACAC CGACACTGTG D T O 524 2701

GGAGAGACAT CCTCTCTGTA Ω ĸ K GTGTAATTCT CACATTAAGA 봈 H 出 GTATGTGCAG CATACACGTC Ø Y V AGCCGGACAC TCGGCCTGTG ٤٠ Ω AACTGCCACA N C H K AGTCCCTGTG TCAGGGACAC H U Ø 544 2761

FIG. 1 CCGAGTGCTA | GGCTCACGAT C 回 GTCTTCCTGG K CAGAAGGACC > CTTTGGAAAG GAAACCTTTC Ċ Ŀ GTGAGGGAGC Ø Ü TK 团 ^begin CGAGAACTGG GCTCTTGACC U J 臼 ĸ GCACGACTTC CGTGCTGAAG r ^ 56**4** 2821

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FIG. 14F TCCTAGGGTG ACGTCGTACT TGGTCTTTGA TGCAGCATGA ACCAGAAACT ATGCAATGAT AAATGCTCCA TTGTGCACCG TTGGGGACTT TTTACGAGGT AACACGTGGC AGGATCCCAC TACGTTACTA AACCCCTGAA Н 臼 . **A** H 工 二 site Σ Ö > Ø Д CATGGGCCAG P L I M CCCCTCATCA GGGGAGTAGT GTACCCGGTC Ø AGGGTCGTGA TTCCGGGACT GAGTGGTTGG Ω CCCGAGAGGG TCCCAGCACT GATCACTTCT X J CTCACCAACC GGCTCTCCC H CTAGTGAAGA H AAGGCCCTGA TK insert Δ, S × 二 J Z Ы H U 0 > 出 O Н S GCTACCCCTG GGAGTCCCGG CATGGACCGG AGCGAATCTG CCCACTCGAC GTACCTGGCC ACACCGACAC GCCGAGCTG CCGGCTCGAC CGATGGGGAC CCTCAGGGCC GGGTGAGCTG TCGCTTAGAC TGTGGCTGTG H ø H Ω Ø H 民 Ш Н U Z 闰 1 Ω U × A K GAGTGTGCGG TGAATAAGTT GCCTGGTTGG ACAAGATGCT Q A K GCCAGGCCAA CGGGTATGGT CGGACCAACC TCCAGAGGGA CGGTCCGGTT AGGTCTCCCT CTCACACGCC ACTTATTCAA GCCCATACCA TGTTCTACGA [24 > Ü U 闰 Σ > о В U Σ Z U Н ¥ AAGTTCTATG H CATGGAGACC 民 D CGGAAGGATT O TTCAAGATAC GGACAGCCAC CCTGTCGGTG CAGATCGCCT ACCAGGAACT TGGTCCTTGA CCGACCAAGG GGCTGGTTCC Ŀı GTACCTCTGG C GCCTTCCTAA GTCTAGCGGA Ω Q P А K Z × × × <u>ប</u> Н K H ပ Ø H E × Д ĸ CGTGTAACAG GGACCGACGG GCACATTGTC ATACATGAAG TATGTACTTC CCTTGTGGAT GGAACACCTA CATTGCCAGT GTAACGGTCA TCTGGACCGG GTTGGAGTCG CCTGGCTGCC AGACCTGGCC CAACCTCAGC Ω ß Y M K > K K L V H Ø H J Ø H Д 644 3061 624 3001 684 704 664 2941 121 181 241 2881 604 584 ന 3 ന SUBSTITUTE SHEET (RULE 26)

ACACCATGCT

GTGGGAGGAC

TTATTACAGG

ACAGCACGGA

AGAGATGTCT

CGGCATGTCC

3301

TGTGGTACGA CACCCTCCTG AATAATGTCC TGTCGTGCCT TCTCTACAGA GCCGTACAGG

回 H H [4 × 民 × Σ H S <u>Ω</u>, Σ 3 Н 744

GTCTCTCACT CAGAGAGTGA TTCAAGTGAT AAGTTCACTA GTACATGGCC CATGTACCGG CTGAAAGCAT GACTTTCGTA ACCTACGGAG TGGATGCCTC GGGGTAAGCG CCCCATTCGC 3361

AGCCATGGTT 3 Д TATGGAAAGC O × U **>**+ GATCTTCACC **[-**נבי Н TCCTCTGGGA 团 3 J H TTCGGGGTGA F G V TGTATGGAGC Ŋ 3 > 764 3421

TCGGTACCAA ATACCTTTCG CTAGAAGTGG AGGAGACCCT AAGCCCCACT ACATACCTCG

ACCTCGCCGG TGGAGCGGCC 回 GGTCGTGTTT H CCAGCACAAA > œ Ü GTAATGGGTT CATTACCCAA Ø H Н TCATTGAGTG AGTAACTCAC U 闰  $\boldsymbol{\vdash}$ AACACGGAGG TTGTGCCTCC NTEV GGTTGAGAGT CCAACTCTCA Ŋ IJ O 3481 784

CCCTTGGTGT GGGAACCACA O Д 回 ACGACCGTCT TGCTGGCAGA ĸ Q 3 U GTACGACCCC CATGCTGGGG Ü J Σ TGTACGATGT ACATGCTACA > Ω CCCAAAGAGG GGGTTTCTCC > ы ም ጽ GGCTCAGACG CCGAGTCTGC R C 804 3541

SUBSTITUTE SHEET (RULE 26)

AGGCCACCCC TCCGGTGGGG H K CGAAACCCCT 봈 GCTTTGGGGA U H K, AATCCTCCAT TTAGGAGGTA 出 H Н AGATCTACAA TCTAGATGTT × >1 Н AACATCAAGG TTGTAGTTCC 团 Н Z GCAGCGGTTG CGTCGCCAAC J ᅂ Ø 3601 824

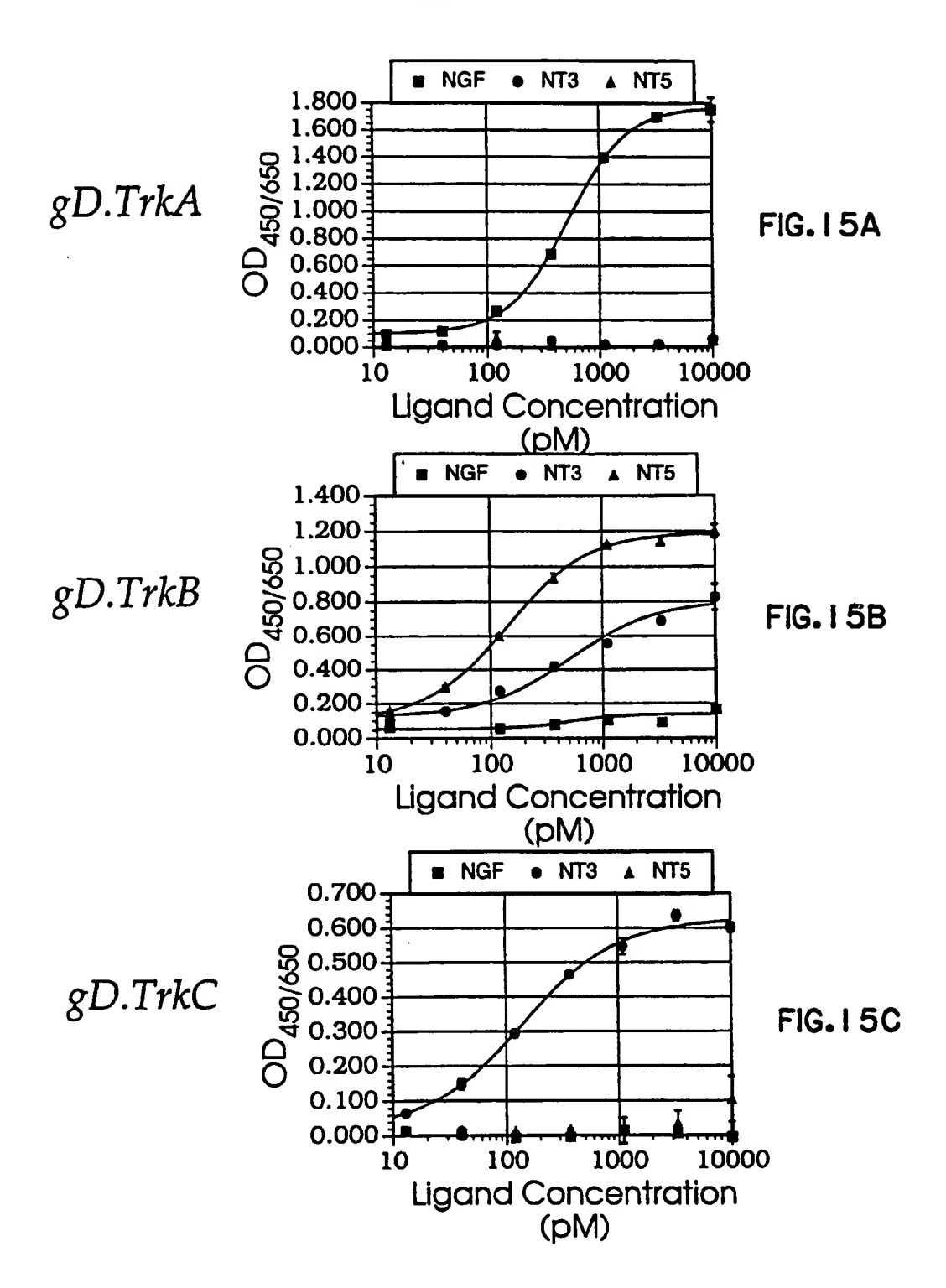
^stop R1 site removed with cut and fill^

844 I Y L D I L G O

TCAATCGATG AGTTAGCTAC ATGAATTAAT TACTTAATT CGACCACCAG GCTGGTGGTC CGATCACCAC GCTAGTGGTG CTGTAAGAAC GACATTCTTG AATCTACCTG TTAGATGGAC 3661

^sv40 early poly A

FIG. 14G GCAATAGCAT CGTTATCGTA TACAAATAAA ATGTTTATT TTATAATGGT AATATTACCA CCCAACTTGT TTATTGCAGC AATAACGTCG GGGTTGAACA GCCCCCATGG CGGCGGTACC 3721



SUBSTITUTE SHEET (RULE 26)

aluI

38/70

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ecoRII
                                                                                                              scrFI
                                                                                                                                               bstNI
                                                                                                                                       dsaV
                                                                                                                                                        apyI
                                                                                                                                                                sexAI
                                                                                                                       mval
                                                                                                                                                                        CAGCAACCAG
                                                                                                                                                                                GTCGTTGGTC
                                                                                      GTCAATCCCA
                                                                              CAGTTAGGGT
                                                                                                                                                                                 GAGTTAATCA
                                                                                                                                                                         CTCAATTAGT
                                                                               GAATGTGTGT
                                                                                       CTTACACACA
                                                                                                                                nsil/avallI
                                                                                                                sfaNI
                                                                               AGAGTCGATC GACAGCTGTG
                                                                                                                                                                         AAGCATGCAT
                                                                                                                                                                                  TTCGTACGTA
                                                                                       CTGTCGACAC
                                                                                                                        ppu10I
                     mbol/ndeII[dam-
                                                                                                                                        nlaIII
                                                               nspBII
                                                                                                                                                                 IHdsu
             IInad
     aluI
                                                       taqI[dam-]
                                                                                                                                                 sphI
                                                                                                                                                         Idsu
                                              dpnII [dam-]
                             dpnI[dam+]
                                      pvuI/bspCI
                                                                       taqI[dam-]
             sau3AI
                                                                                        TCTCAGCTAG
                                                                                                                                                                           GAAGTATGCA
                                                                                                                                                                                  CTTCATACGT
                                                                mcrI
                                                                                                                                                                                              FIG. 1 6A
                                                       hinfI
                                              pleI
                                                                         mael
                                                                 rmaI
                                                                                 ATTATTGACT
                                                                                                                                                                           CCAGCAGGCA
                                                                                         TAATAACTGA
                                                                                                                                                                                   GGTCGTCCGT
                                                                                                                                                            apyI[dcm+]
                                                                                  CCCGACATIG
                                                                                                                                                                                    GGGTCCGAGG
                                                                                          GGGCTGTAAC
                                                                                                                                                                            CCCAGGCTCC
                                                                                                          nlaIV
                                                                                                                                  ecoRII
                                                                                                                  scrFI
                                                                                                                                                   bstNI
                                                                                                                                           dsaV
                                                                                                                                                                    bsaJI
                                                                                                                           mvaI
                        hgiAI/aspHI
                               ec1136II
                                                                                  TTCGAGCTCG
                                                                                                                                                                             GTGGAAAGTC
                                                                                                                                                                                    CACCTTTCAG
                                                                                           AAGCTCGAGC
                                        bsp1286
                                                 bsiHKAI
                hgiJII
                                                                 banII
                                                          bmyI
        sacI
sstl
                                                                           taqI
                                                                                                                                                                             71
```

mnlI

AGAAGTAGTG

TCTTCATCAC

CTCGATAAGG

GAGCCGGAGA

GGCTCCGGCG

ATACGTCTCC

AAAAAAATAA

CGACTGATTA

GCTGACTAAT

281

TTTTTTTT

TATGCAGAGG

CCGAGGCCGC

bsaJI acil

mnll

CTCGGCCTCT

GAGCTATTCC

aluI

mnlI

bsaJI

haeIII/palI

mnll

mnlI

ddeI

haeIII/palI

39/70 nlaIII GTCAGCAACC GGCGGGGTAC CCGCCCCATG bsaJI styI ncol dsaI bsliacil ATCTCAATTA GCGGGTAAGA TAGAGTTAAT CGCCCATTCT acil ppul0I GGGGATTGAG GCGGGTCAAG GTTTCGTACG CGCCCAGTTC CAAAGCATGC nspHI nspI sphI bsrI nsil/avaIII haeIII/palI aciI sfaNI fnu4HI CCCCTAACTC CAGAAGTATG GTCTTCATAC nlaIII pdlI sfiI acil GCCCATCCCG CGGGTAGGGC CCCCAGCAGG GGGGTCGTCC fokI apyI[dcm+] aciI nlaIV CCCTAACTCC GGGATTGAGG AGGGGTCCGA TCCCCAGGCT bstNI bsaJI mval ecoRII SCrFI dsaV ATAGTCCCGC CACACCTTTC acil GTGTGGAAAG [dcm+] 211 141

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FIG. 16B

CCAAAATATG TAGAGCGATA ATCTCGCTAT GGTTTTATAC pflMI bslI /xmaIII/eclXI scfI GTACCGCCTA CATGGCGGAT TCGCCGTGTC AGCGGCACAG haeIII/palI acil csp6I donor rsaI eagI/ mcrI eaeI hpall cfrI Idsm GCTTATCCGG CGAATAGGCC AACTGCATCG TTGACGTAGC AGTGACGTAA TCACTGCATT ^splice sfani maeII maeIII aluI rmaI maeI nheI CAAAAAGCTA CCGTGCCAAG GTTTTTCGAT AGCTGGTAAC GGCACGGTTC TCGACCATTG aluI FIG. 1 6C taqI fnuDII/mvnI TAGGCTTTTG ATCCGAAAAC CGCGGATTCC GCGCCTAAGG CATGGT nlaIII GGTAGTACCA hinFI tfiI 1236I IT DHFR ATG^ bsh1 thaI bstU CCAT aci haeIII/pa rmaI bsaJI avrll maeI styl blnI fnu4HI AAACCTCCGG TAGGGGGGAC TTTGGAGGCC ATCCCCGCTG nspBII TGCATTGGAA **PPAI** ACGTAACCTT stuI haeI mnlI acil TCCTCCGAAA GGCCCTTGCC TCTCCTAAAA AGGAGGCTTT CCGGGAACGG AGAGGATTTT caull mnlI hpaII SCrFI dsaV Idsm ncil 351 401 471 SUBSTITUTE SHEET (RULE 26)

mnlI

bslI

41/70

ddeI

apyI[dcm+]

TCTCCATTCC

AGAGGTAAGG

CAAAGAATGA GTTTCTTACT GTTCATGAAG CAAGTACTTC csp6I rsal scal CCTTGCTCAA GGAACGAGTT asp700 xmnI ddeI CCTCCGCTCA GGAGGCGAGT bsrBI haeIII/palI acil mnlI apyI[dcm+] haeI ecoRII CCTACCCTGG GGATGGGACC SCLFI bstNI dsaV mvaI bsaJI bsmAI GGGATTGGCA AGAACGGAGA bsal TCTTGCCTCT CCCTAACCGT

ecoRII SCrFI bstNI dsaV mval eco57I

TTTGGACCA AAAACCTGGT sexAI TATGGGTAGG ATACCCATCC GGTAAACAGA ATCTGGTGAT CCATTTGTCT TAGACCACTA hphI alwNI TTCAGTGGAA AAGTCACCTT GGTGTTGGAG CCACAACCTC mnlI

hinfI

earI/ksp632I

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Ilodm

541

ddeI aseI/asnI/vspI tru9I msel ahaIII/draI tru9I mseI mboll taql hinfI tfiI

ACCACCACGA TGGTGGTGCT AACTCAAAGA TTGAGTTTCT CTCAGTAGAG GAGTCATCTC ATTATATCAA AGGACAGAAT TAATATAGTT TCCTGTCTTA CGACCTTTAA GCTGGAAATT TGAGAAGAAT ACTCTTCTTA 681

3.16D FIG

hgiAI/aspHI

hgiJII

sstI

ec1136II

**bsp1286** 

**bsiHKAI** 

banII

bmyI

aluI

751

42/70 haeIII/palI ddeI ATGAATCAAC CAGGCCACCT TACTTAGTTG GTCCGGTGGA TIGGCAAGTA AACCGTTCAT apyI[dcm+] haeI bstNI mvaI ACAACCGGAA TGTTGGCCTT hpall ecoRII Idsm bsaWI scrFI dsaV hinfI nlaIII tfiI CCAGGAAGCC GACTTATTGA CTGAATAACT apyI[dcm+] ecoRII SCLFI bstNI dsaV mvaI aflII/bfrI tru9I GATGCCTTAA CTACGGAATT sfaNI mseI GGAT fokI TTCAAACCTA mnlI AAGTTT bstXI AAGAACGGTT TTCTTGCCAA accI nlaIII CCTCGAGTAA GGAGCTCATT

FIG. 1 6E

GGTCCTTCGG

GTTCTGTTTA

GTCGGAGGCA

AAGTAGACAT GGTTTGGATA

821

CAGCCTCCGT

CCAAACCTAT

TTCATCTGTA

CAAGACAAAT

ų.

TGATTTGGGG ACTAAACCCC AGGGTCTTTA TCCCAGAAAT CTGTGCAAAA GACACGTTTT maeII aflIII maeIII GTGACAAGGA TCATGCAGGA ATTTGAAAGT TAAACTTTCA mboI/ndeII[dam-]
dpnI[dam+] apol AGTACGTCCT dpnII[dam-] alwI[dam-] nlaI sau3AI CACTGTTCCT maeIII TAGACTCTTT ATCTGAGAAA hinfI pleI 891

II

mnlI bslI ddeI GTCCTCTG CAGGAGAGAC ecoNI mnlI apyI[dcm+] bsaJI bs TATGGGTCCG ATACCCAGGC ecoRII dsaV bstNI SCLFI mval

bs

AAATATAAAC CTCTCCCAGA TTTATATTTG GAGAGGGTCT

961

mnlI

ahaII/bsaHI

hinlI/acyI

hgaI

FIG. 1 6F

sfaNI apyI[dcm+] ecoRII bstNI mvaI dsaV sau96I avall

SCLFI

TCCTTCTACG AGGAAGATGC mbolI DHFR ^END **AAAGACTAAC** TTTCTGATTG mbolI CTACGAGAAG GATGCTCTTC accI TCAAACTTCA AGTTTGAAGT GGAAAAGGC ATCAAGTATA CCTTTTTCCG TAGTTCATAT sfaNI mnlI AGGTCCAGGA TCCAGGTCCT asuI 1001

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nlaIII styI ncoI ppul0I dsaI

mnli alui nsii/avaiii bsaJi CCCC TCCTAAAGCT ATGCATTTTT ATAAGACCAT GC

GGGACTTTG CCCTGAAAAC TATTCTGGTA TACGTAAAAA AGACGAGGGG AGGATTTCGA TCTGCTCCCC TCCTAAAGCT AAAGTTCAAG TTTCAAGTTC

1071

FIG. 16G

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apyI[dcm+] sau96I avall asuI ATAGTATGTG TATCATACAC CTCCCAGGTC GAGGGTCCAG ecoRII SCrFI bstNI dsaV bsaJI mvaI bslI ATTGGAATAC TAACCTTATG CAGGTGTCCA GTCCACAGGT aseI/asnI/vspI AATTAATACA TTAATTATGT TTTCTCCCA AAAGAGAGGT tru9I mseI fnuDII/mvnI CCACTTTGCC TGCGCCGATG ACGCGGCTAC **bsh1236I** fnu4HI bstUI acil thal foki TAGATAACAT ATCTATTGTA CTTCGTTAGA GAAGCAATCT mbol/ndeII[dam-] scfI TAGGGGAACC GGTGACACTA CCACTGTGAT CTGGCTTTAG ATCCCCTTGG bsaJI dpnII[dambstYI/xhoII dpnI[dam+] alwI[dammaeIII hphI sau3AI GACCGAAATC ATACGATTTA TATGCTAAAT 1131 1201 SUBSTITUTE SHEET (RULE 26)

45/70

FIG. 16H

FIG. 1 61

1271

46/70

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mpoI/ndeII[dam-]
                                    xmal/pspAI
                                                                                                                             caull
                                                                                                                                            bsaJI
                                                                                                                      dpnII[dam-
                      hpall
SCLFI
                                                                                                                                    bstYI/xhoII
                                                                                                              dpnI[dam+]
                                                                                                                                                          alwI[dam-
                                                                         caulI
                                                                                 bsaJI
      nciI
              Idsm
                                                   scrFI
                             dsaV
                                                                                                                                                   alwI[dam-]
                                                                  dsaV
                                           smal
                                                           ncil
                                                                                         aval
                                                                                                                                                                  mnll bsaJI
                                                                                                                                                                         GAGGATCCCC
                                                                                                                                                                                CTCCTAGGGG
                                                                                               sau3AI
                                                                                                                                            bamHI
                                                                                                                              nlaIV
                                                                                                                                                          hincII/hindII
                                                                                                                                                   maeI
                                                                                                                                             rmaI
                                                                                                                                                                  xbaI
                                                                                                                                                                                 CAGCTGAGAT
                                                                                                                                                                          GTCGACTCTA
                                                                                                                              pleI
hinfI
                                                                                                                                            taqI
salI
                                                                                                                                                                  accI
                                                                                                                                                            ndIII bspMI
bsgI
GCTTCTGCAG
CGAAGACGTC
                                                                                                                                                     pstI
                                                                                                                                                            hindIII
                                                                                                                                              scfI
aluI
                                                                                                                                                                    ddeI
                                                                                                                                                                           TCGGTTCTAA
                                                                                                                                                                                  AGCCAAGATT
                                                                                                                                                                    bsaJI
                                                                                                                                                             mnlI
                                                                                                                                                                           CAACTGCACC
                                                                                                                                                                                  GTTGACGTGG
```

AAATAAAGCA TTTATTTCGT maeIII ATTACCAATG TAATGGTTAC TTGCAGCTTA AACGTCGAAT fnu4HI aluI Indd AACTTGTTTA TTGAACAAAT ^sv40 sau96I haeIII/palI 6 bsaJI GCCATGGCCC CGGTACCGGG asuI nlaIII taqI haeIII/palI styI ncoI dsaI fnu4HI acil clai/bsp106 pall ATCGATGGCC TAGCTACCGG sfil cfrI eaeI GGGGAATTCA CCCCTTAAGT ecoRI apol 1321

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GGTTTGAGTA CCAAACTCAT ACACCAAACA TGTGGTTTGT CGTAAGATCA GCATTCTAGT maeI bsmI TTATTTCGTA AAAAAGTGA TTTTTCACT AATAAAGCAT AAATTTCACA TTTAAAGTGT apol TATCGTAGTG ATAGCATCAC sfaNI

1391

rmaI

FIG. 16J

```
GTACCTTCTG AGGCGGAAAG
                                                                                                                                                                                                             TCCGCCTTTC
                                                                                                                                                                                              acil
                                                                                                                                                                                      mnlI
                                                                                                                                                                                              ddeI
                                                                                                                                                                                                              CATGGAAGAC
                                                                                                                                               csp61
                                                                                                                                       rsal
                                                                                                                                                                                      asp718
                                                                                                                                                                                              acc651
                                                                                                                                                                       hgiCI
                                                                                                                                                       nlaIV
                                                                                                                                                                               banI
                                                                                                                                                               kpnI
                                                                                                                                                                                                       ACTIGGTIAG
                                                                                                                                                                                                              TGAACCAATC
                                                                                        aseI/asnI/vspI
                                                                                                                                                                                                       TGAAAGAGGA
                                                                                                                                                                                                               ACTTTCTCT
                                                                                                        GGAATTAATT
                                                                                                               CCTTAATTAA
                                                                                                                                                                                               mnlI
                                                tru9I
        mbol/ndeII[dam-
                                                                mseI
                                                       claI/bsp106[dam-]
                                                                                              asp700
                                                                        mboI/ndeII[dam-]
                                                                                 xmnI
                        dpnII[dam-
               dpnI[dam+]
                               pvul/bspCI
                                                                                                                        origin^
                                              taq1[dam-]
                                                                               dpnI[dam+] >
dpnI[dam+]
sau3AI
                                                                                                                CCTAGCTAGC
                                                                                                        GGATCGATCG
                                                                                                                                                                                                       AAATAACCTC
                                                                                                                                                                                                               TTTATTGGAG
                                                                                                                                                                                               mnlI
                                                                                                                                                                                                                          FIG. I 6K
                                                                                                alwI[dam-]
                                      mcrI
                                                                sau3AI
                                                                                                                       sv40
                                                                                                                                                H
                                                                                                                                                haeIII/pal
                                                                                                                                                                                                       CCATGGCCTG
                                                                                                                                                                                                               GGTACCGGAC
                                                                                                        TATCATGTCT
                                                                                                                ATAGTACAGA
                                                                                                 nlaIII
                                                                                                                                                       haeI
                                                                                                                                                                                               nlalII
                                                                                                                                                                                       bsaJI
                                                                                                                                                                               dsaI
                                                                                                                                                                styl
                                                                                                                                                                       ncol
                                                                                                                                                                                               hhaI/cfoI
                                                                                                                                                                                                       CGGCGCAGCA
                                                                                                        CAATGTATCT
                                                                                                               GTTACATAGA
                                                                                                                                                                                                               GCCGCGTCGT
                                                                                                                                                                        fnu4HI
                                                                                                                                                                               Ivqq
                                                                                                                                                                                       hinPI
                                                                                                         1461
                                                                                                                                                                                                        1501
                                                                    SUBSTITUTE SHEET (RULE 26)
```

	AGAAGTATGC	GCAGAAGTAT	acii cc GCCCCTAACT
	n+] c cccagcaggc g gggrcgrccg	nlarv I NI 'I[dcm+] 'I GGC TCCCCAGCAG	ciI fokI cgcccaTC
nlalv scrfI mvaI	ecoRII dsaV bstNI apyI[dcm+] bsaJI CCCCAGGCTC C	mva bst bsad GTCCCCA	I CCCCTAACT GGGGATTGA
	TGTGGAAAGT	scrfi mval ecoRII dsaV bstNI apyl[dcm+] exAI ccA GGTGTGGAAA GGT CCACACCTTT	acil CATAGTCCCG
	TCAGTTAGGG	scrfi mva] ecoRII dsaV bsth apy] rcAGCAACCA AGTCGTTGGT	AGTCAGCAAC
	GGAATGTGTG	TCTCAATTAG	sfaNI u101 iI/avaIII III G CATCTCAATT G CATCTCAATT C GTAGAGTTAA
FIG. 1 6L	aluI pvuII nspBII AACCAGCTGT TTGGTCGACA	sfaNI ppul0I nsil/avaIII nlaIII sphI nspI nspHI nspHI TTTCGTACGT	sfa ppul0 nsil/ nlaIII sphI nsplI nspHI gcaaagcarg c
	1571	SUBSTITUTE SHEET (RU	JLE 26)

haeIII/palI avrII bsaJI blnI mnlI TTTTGGAGGC AAAACCTCCG TTATGCAGAG AATACGTCTC haeI stuI mnlI GAGGAGGCTT AAAAAAATA TTTTTTTT mnlI mnlI CCGACTGATT GGCTGACTAA GTCTTCATCA CAGAAGTAGT nlaIII bsaJI TCCGCCCCAT ACTCGATAAG TGAGCTATTC styI ncol dsal aluI acil bslI haeIII/palI ddeI GGCGGGTAAG CCTCGGCCTC CCGCCCATTC bsaJI mnlI haeIII/palI mnlI mnlI mnlI acil fnu4HI bsaJI aciI bglI sfiI GCCGAGGCCG CGGCTCCGGC GGGGGGTCAA CCGCCCAGTT haeIII/palI bsrI asil 1841 1781

FIG. I 6M

FIG. 16N

ATTGTCGAAC

```
CCAACTTAAT
                                                                                                                                                                                     GGTTGAATTA
                                                                              ATCCTGCAGG
                              bspMI
                                                                      sse8387I
                                                                                      TAGGACGICC
                                                                                                                                                              tru9I
                                                                                                                                                                     mseI
                                                              psdl
                                             pstI
                                      scfI
                                                      ahaIII/draI
                                                                                                      site
                                                                                                                                                                      maeIII
                                               tru9I
                                                                                                                                                                                     GACCGCAATG
                                                                                                                                                                              CIGGCGTTAC
                                                              mseI
                                                                               CGCCATTTAA
                                                                                      GCGGTAAATT
                                                                                                       Hpa1
                                                                      swal
                                                                                                                                                              apyI[dcm+]
                      fnuDII/mvnI
       hhaI/cfoI
                                                              bsh1236I
                                                                                                       inserted into
                                               hhal/cfol
                                                                                                                                       ecoRII
                                                                                                                                                       bstNI
                                                                                                                       SCRFI
hinPI
                                                                                                                                               dsaV
                                                                       msel bssHII
                                                                                                                               mvaI
                                                                                                                                                                      bsaJI
                               bstuI
                                      hinPI
               thaI
                                                                               GCGGCCGCTT AATTAAGGCG
CGCCGGCGAA TTAATTCCGC
                                                                                                                                                                                      ACCCTTTTGG
                                                                                                                                                                               TGGGAAAACC
                                                       ascI
                                                               tru9I
                      eagI/xmaIII/eclXI
                                                                                                        'linearization linker
                                               tru91
       haeIII/palI
                                                                                                                                                                        bsrI
                                                              mseI
                                                       pacI
                                                                                                                                                               maeIII
                                                                                                                                                                               ACGTCGTGAC
                                                                                                                                                                                      TGCAGCACTG
                                                                       bsrBI fnu4HI
aciI
                                                        fnu4HI
                                               cfrI
                                                                                                                                                                       maell
                               eaeI
                mcrI
                                                                aciI
                                       notI
                                                paeR7I
                                                                                                                                                                                       AGCAAAATGT
                                                                                        CAATGGAGCT
                                                                                                                                                                               TCGTTTTACA
                                                                                 GTTACCTCGA
                                taqI
                                                        avaI
                                        xhoI
                                                                mnlI
                                                                                                start pUC118
                                                                        maeIII
                                                                                                                                                haeIII/palI
                                                                                                                                                                                GCACTGGCCG
                                                                                                                                                                                        CGTGACCGGC
                                                                         aluI
                                                                                 GCAAAAAGCT
                                                                                         CGTTTTCGA
                                                                                                                                                         eaeI
                                                                                                                                                                cfrI
                                                                                                                                                                       bsrI
                                                                                         GATCCGAAAA
                                                                                                                                                                                 TAACAGCTTG
                                                                                 CTAGGCTTTT
                                                                                                                                                                         aluī
                                                                  rmaI
                                                                         maeI
                                                                                                                                                                 maeIII
                                                                                                                                                                                 1971
                                                             SUBSTITUTE SHEET (RULE 26)
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```
mbol/ndell[dam-]
                                                                                                                                                                                  CTTACGCATC
GAATGCGTAG
                                                                                                                                                                          sfaNI
                                   dpnII[dam-
                                             pvuI/bspCI
                           dpnI[dam+]
                                                                                                                                                                                           CATAAAAGAG
                                                                                                                                                                                   GTATTTCTC
                                                      mcrI
                                                               CCGCACCGAT
                                                                        GGCGTGGCTA
sau3AI
                                            mboll acil
earl/ksp6321
         sau96I
                  haeIII/palI
                                                                                                                                                         acil
                                                                        CGCTTCTCCG
                                                                                                                                                                                   GCCTGATGCG
                                                               GCGAAGAGGĈ
                                                                                                                                                                                            CGGACTACGC
                                                                                                                                                                  sfaNI
                                    mnlI
                                                                                                                                                                           ahall/bsaHI
                                                                                                    hhaI/cfoI
                                                                                                                                       hinlI/acyI
                           asuI
                                                                                           hinPI
                                                                                                             nlaIV
                                                                                                                                                hgiCI
                                                                                                                                                         haeII
                                                                                                                                                                  banI
                                                                                                                       narI
                                                                                                                               kasI
                                                                                                                                                                                           CCGCTTACCG
                                                                                                                                                                                    GGCGAATGGC
                                                                         ACCGCATTAT
                                                                TGGCGTAATA
                                                       nspBII
                                              pvull
                                      aluI
                                                                CTTCGCCAGC
                                                                         GAAGCGGTCG
                                                                                                                                                                                     TAGCCTGAAT
                                                                                                                                                                                             ATCGGACTTA
                                                                                                                                                                             pal
                                                                          GTGTAGGGGG
                                                                                                                                                                                      AACAGTTGCG
                                                                                                                                                                                             TTGTCAACGC
                                                                  CACATCCCCC
                                                         fokI
                                                fru4HI
                                                                                                                                                                                     CGCCCTTCCC
                                                                                                                                                                                              GCGGGAAGGG
                                                         bbvI
                                                                           GCGGAACGTC
                                                                  CGCCTTGCAG
                                                                                                                                                                                      2101
                                                                  2041
```

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FIG. 160

hhaI/cfoI

hinPI

hhaI/cfoI hinPI fnuDII/mvnI scfI bsh1236I bstuI

hinPI

thal

fnu4HI aciI hhaI/cfoI rsaI

bslI csp6I

CATCGCCGCG GTAGCGGCGC TACGCGCCCT ATGCGCGGGA CGTTGGTATC GCAACCATAG TATGCAGTTT ATACGTCAAA maeII

fnu4HI hinPI fnu4HI

AAGTGTGGCG

ACACGCCATA

TGTGCGGTAT

2171

acil

TTCACACCGC

aciI

fnuDII/mvnI bctUI thaI

SUBSTITUTE SHEET (RULE 26)

acil hhaI/cfoI hinPI

hhaI/cfoI

fnuDII/mvnI bsh1236I bstUI

maeIII bbvI

msel bsh1236I

ATTAAGCGCG

2231

TAATTCGCGC

aciI

tru9I

GCTACACTTG CGATGTGAAC aciI CAGCGTGACC GTCGCACTGG maeIII TGGTTACGCG ACCAATGCGC GCGGGTGTGG CGCCCACACC

hpall Idsm

bsp1286

banII

bmyI

hgiJII

nlaIV

AGCGCCCGCT

CCAGCGCCCT

GGTCGCGGGA

bsrBI

hhaI/cfoI

haeII

hinPI

rmaI

acil

maeI

haeII

TCGCGGGCGA

naeI

AATCGGGGGC TTAGCCCCCG TCAAGCTCTA AGTTCGAGAT aluI GCTTTCCCCG CGAAAGGGGC cfr10I ACGITCGCCG TGCAAGCGGC maeII CTTTCTCGCC GAAAGAGCGG

TCTTCCCTTC

CCTTTCGCTT

2301

GGAAAGCGAA

Ilodm

AGAAGGGAAG

. 16P

FIG

maelI

hinfI

pleI

mnll

hgiCI nlaIV

taqI banI

Ihqh CTTGATTTGG CCCCAAAAA GGCACCTCGA

GCTTTAC

AGGGAAATCC CAAGGCTAAA

GTTCCGATTT

TCCCTTTAGG

2371

nlaIV

GAACTAAACC GGGGTTTTT CCGTGGAGCT CGAAATG AGT TCA

TIGACGIIGG AGICCACGII maeII drdI TTTTCGCCCT GATAGACGGT CCATCGCCCT /pall haeIII sau96I asuI ACGTAGTGGG maeII draIII bsaAI

TCAGGTGCAA AACTGCAACC AAAAGCGGGA CTATCTGCCA GGTAGCGGGA TGCATCACCC GTGATGGTTC CACTACCAAG

2401

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TTTTGATTTA AAAACTAAAT CGGGCTATTC aval AACCCTATCT bslI bslI AACAACACTC TCCAAACTGG bsrI GGACTCTTGT hinfI pleI CTTTAATAGT tru9I mseI

GCCCGATAAG TTGGGATAGA TTGTTGTGAG AGGTTTGACC CCTGAGAACA GAAATTATCA 2501

tru9I tru9I

tru9I

mseI

ACAAAAATTT apol mseI AGCTGATTTA aluI TTAAAAATG msel haeIII/palI GGCCTATTGG TGCCGATTTC

TGTTTTAAA TCGACTAAAT AATTTTTAC CCGGATAACC ACGCCTAAAG TAAGGGATTT ATTCCCTAAA 2571

FIG. 1 60

	55/70	
rsal rsal cagracaatc Grcatgrag	hinPI fnu4HI nlaIII hhaI/cfoI /aspI bbvI GTCATGGCTG CGCCCGACA CAGTACCGAC GCGGGGCTGT	maeIII aciI ccGCTTACAG ACAAGCTGTG GGCGAATGTC TGTTCGACAC
hgiAI/aspH bsp1286 bsiHKAI bmyI dde apaLI/snoI alw44I/sno GGTGCACTCT CCACGTGAGA	bsrI maeIII saAI tth111I ACGTGACTGG TGCACTGACC	mspl hpall scrfi ncil dsaV fokI caull CTCCCGGCAT
maeli psp14061 tru91 msel ATTAACGTTT ACAATTATA	bsrI maeIII acil maeII ccGCATAGTT AAGCCAACTC CGCTATCGCT ACGTGACTGG GGCGTATCAA TTCGGTTGAG GCGATAGCGA TGCACTGACC hinPI	hhal/cfol thal fnuDII/mvnI bstUI bsh1236I al drdi cG CGCCCTGACG GGCTTGTCTG GC GCGGACTGC CCGAACAGAC
vnI tru9I msel sspI TTAACAAAAT AATTGTTTTA	nu4HI tru I aciI mse CCGCATAGTT GGCGTATCAA	hh tha fnu bst nspBII bsh aciI hgaI cccGCTGACG GGGCGACTGC
thal fnuDII/mvnI apol bstUI tr bsh1236I ms TTGCGCAATT TT	sfan 2691 TGCTCTGATG ACGAGACTAC	acil 2761 CCCGCCAACA GGGCGGTTGT

FIG. 16S

fnuDII/mvnI

bsh1236I

ahaII/bsaHI

aatII

hinII/acyI

hinPI

bstUI

56/70

CAGTATTCTT GTCATAAGAA fnuDII/mvnI fnuDII/mvnI bsh1236I hhaI/cfoI ACGCGCGAGG TGCGCGCTCC thal mull bsh1236I bstuI thal hinPI bstuI acil thaI nlallI IHdsq CATCACCGAA TAATGTCATG GTAGTGGCTT ATTACAGTAC rcal hphI tru9I msel TTTTATAGGT AAAATATCCA AAAAGTGGCA TTTTCACCGT hphI mnlI GTGTCAGAGG CACAGTCTCC GGGCCTCGTG ATACGCCTAT aluI nlaIII haeIII/palI eco01091/draII nspHI GGAGCTGCAT CCTCGACGTA nspI fnu4HI Inqq mnlI sau96I asuI hpaII caull SCrFI Idsm dsaV nciI ACCGTCTCCG GAAGACGAAA CTTCTGCTTT bsmAI esp3I bslI bpuAI Ilodm ppsI 2831 2901

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hhaI/cfoI GAAATGTGCG CTTTACACGC ACTTTTCGGG TGAAAAGCCC GTCAGGTGGC ddeI maeII AAAGAATCTG TTTCTTAGAC ATAATAATGG TATTATTACC 2951

bsmAI

rcal

nlaIII

IHdsq bsrBI aciI

ATAACCCTGA TATTGGGACT TCATGAGACA AGTACTCTGT TACATAGGCG ATGTATCCGC ACATTCAAAT TGTAAGTTTA GATTTA CTAAAT TTTT AAAA

TAAACAAATA

GCCTTGGGGA

3071

CGGAACCCCT

3001

nlaIV

ATTTGTTAT

Ilodm

earI/ksp632I

CTTATTCCCT GAATAAGGGA CCGTGTCGCC TICAACATIT AGTATGAGTA AAAAAGGAAG AATAATAG sspI

GGCACAGCGG AAGTTGTAAA TCATACTCAT TTTTCCTTC TTATTATAAC TAAATGCTTC

ATTTACGAAG

sfaNI AAAGTAAAAG AACGCTGGTG hphI hphI fnu4HI acil

TTTCATTTC TTGCGACCAC CTCACCCAGA GAGTGGGTCT CCTGTTTTG GGACAAAAAC TAAAACGGAA ATTTGCCTT AAAAACGCCG TTTTGCGGC

3141

SUBSTITUTE SHEET (RULE 26)

hgiAI/aspHI

**bsp1286** 

mpol/udeII[dam-]

sau3AI

dpnII[dam-]

dpnI[dam+]

bstYI/xhoII

acil

alwI[dam-]

nspBII

bsiHKA1 mpoI/ndeII[dam-] sau3AI

bmyI dpn1[dam+]

apall/snol dpnII[dam-] mboll[dam-]

bsrI taql maeIII lous, alw44I/ eco57I

AACAGCGGTA TTGTCGCCAT TGACCTAGAG ACTGGATCTC GTTACATCGA CAATGTAGCT ATGCTGAAGA TCAGTTGGGT GCACGAGTGG CGTGCTCACC AGTCAACCCA TACGACTICT 3201

**6**1 <u>FIG.</u>

FIG. I 6U

sau3AI

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fokI ahaIII/draI TTTAAAGTTC AAATTTCAAG CATCTTACGG GGTCGCCGCA GTAGAATGCC CCAGCGGCGT fnu4HI acil tru9I mseI sfaNI hgiAI/aspHI MCLI GTGTCTTTTC TCTCGTTGAG CACAGAAAAG AGAGCAACTC GATGAGCACT CTACTCGTGA **bsp1286 bsiHKAI** bmyI bcgI maeIII TGAGTGGTCA ACTCACCAGT CAAAAGGTTA TGCGGCCCGT GTTTTCCAAT ACGCCGGGCA bsrI ahaII/bsaHI SCLFI hpall caulI hinl1/acyI ncil Idsm dsaV Ihqh psp1406I hgaI csp6I scal 1 maeII asp700 rsal TCCCGTGATG AGGCACTAC TTGGTTGAGT GGGCTTCTTG AACCAACTCA CCCGAAGAAC xmnI Ilodm fnuDII/mvnI GAGTTTTCGC CTCAAAAGCG CGCGGTATTA TACACTATIC TCAGAAIGAC AIGIGATAAG AGICTIACIG **bsh1236I** hinPI hhaI/cfoI mboI/ndeII[dam-] bstui acil thaI ddeI dpnII[dambstYI/xhoII dpnI[dam+] alwI[dam-] TGCTATGTGG ACGATACACC AGATCCTTGA TCTAGGAACT 3321 3381 3261

fnu4HI

59/70

mbol/ndeII[dam-

dpnII[dam-]

alwI[dam-]

nlaIII

dpnI[dam+]

sau3AI maeIII

nlaIII

GGTTGAATGA CCAACTTACT haeIII/palI cfrI eaeI fnu4HI AACACTGCGG aciI GTACTCACTA CATGAGTGAT nlaIII GACGGTATTG CTGCCATAAC fnu4HI **I**vqq TTATGCAGTG AATACGTCAC TCATTCTCTT AGTAAGAGAA TACCGTACTG ATGGCATGAC nlaIII

sau96I

avaII asuI sau3AI SUBSTITUTE SHEET (RULE 26)

3441

mbol/ndell[dam-] dpnI[dam+]

dpnII[dampvuI/bspCI

aluI CGAAGGAGCT mnlI mcrI

TIGCACAACA IGGGGGATCA IGTAACICGC ACATTGAGCG ACCCCCTAGT AACGTGTTGT AACCGCTTTT TTGGCGAAAA acil GCTTCCTCGA TCTGACAACG ATCGGAGGAC AGACTGTTGC TAGCCTCCTG 3511

Idsm

aluI mbol/ndeII[dam-] nlaIV sau3AI

hpall dpnII[dam-] bsaWI dpnI[dam+]

Inqq ATGCCAGCAG TACGGTCGTC sfaNI ACGACGAGCG TGACACCACG ACTGTGGTGC maeIII TGCTGCTCGC GCCATACCAA CGGTATGGTT GCTGAATGAA CGACTTACTT CTTGATCGTT GGGAACCGGA GAACTAGCAA CCCTTGGCCT 3581

FIG. 1 6V

FIG. 1 6W

60/70

												H	CTGG	GACC							
										palI	Idsm	hpall	Trccggcrgg	AAGGCCGACC							
Idsm	hpail	SCLFI	aluI nciI	rmaI dsaV	mael caull	ACTIACTCIA GCTICCCGGC	TGAATGAGAT CGAAGGGCCG	Ilgd	sau96I	haeIII/palI	hinPI asuI	hhaI/cfoI	CGCTCGGCCC	TGGTGAAGAC GCGAGCCGGG A	thaI	fnuDII/mvnI	bstUI		$\vdash$		AGAGCGCCAT AGTAACGTCG
			bsrI	<b>—</b>		CTGGCGAACT	GACCGCTTGA			sau96I	avall	Inse	AAGTTGCAGG	TTCAACGTCC		-		bsmAI	bsaI	TGAGCGTGGG	ACTCGCACCC
	hhaI/cfoI			tru9I	mseI	AAACTATTAA	TTTGATAATT				acil	mpli	GAGGCGGATA	CTCCGCCTAT	Idsm	Iledu	cfr101	nlaIV hphI	<pre>lmdq/Insb</pre>	CTGGAGCCGG	GACCTCGGCC
hinPI	hhaI,	mstI	aviII/fspI	maeII	psp1406I	AACGTTGCGC	TTGCAACGCG			fokI	bsrI	n	•	TCTGACCTAC						GCTGATAAAT	CGACTATTTA
						CAATGGCAAC	GTTACCGTTG			tru9I	mseI	se/Tese	AACAATTAAT	TTGTTAATTA						CTGGTTTATT	GACCAAATAA
						3651							3711							3781	
								SUB	STI	TU	FE S	SH	EET	(RUL	E 26	)					

h

TGAGTTTTCG

TCCCTTAACG

TGATAATCTC ATGACCAAAA

ATTTAAAAGG ATCTAGGTGA AGATCCTTTT

maeI

ahaIII/draI

TAGATCCACT

TAAATTTTCC

4021

**LHqsd** 

rcal

bstYI/xhoII

mboll[dam-]

msel

tru9I

AGGGAATTGC

TACTGGTTTT

ACTATTAGAG

TCTAGGAAAA

61/70

FIG. 16X tru9I maeII msel GGAGTCAGGC TTAAGCATTG TTCATTTTA AATTCGTAAC AAGTAAAAT hinfI pleI tru9I mseI eam1105I ahaIII/draI CTAAATTTTG TACACGACGG ATGTGCTGCC GCCTCACTGA CGGAGTGACT GATTTAAAAC tru9I mseI mnlI nlaIII hgiCI nlaIV banI TGAGATAGGT ACTCTATCCA CGTAGTTATC GCATCAATAG TGAAATCTAA ACTITAGATI mboI/ndeII[dam-] mbol/ndeII[dam-] dpnII[dam-] dpnI[dam+] ddeI dpnII[damalwI[dam-] dpnI[dam+] CTGTCTAGCG GACAGATCGC GGAGGGCATA TGAGTATATA ACTCATATAT CCTCCCGTAT sau3AI sau3AI mnll mbol/ndeII[dam-] GACCAAGTTT GAACGAAATA CTTGCTTTAT GATGGTAAGC CTACCATTCG CTGGTTCAAA sau3AI hphI dpnII [dambstYI/xhoII dpnI[dam+] alwI[dam-] rmal bsrI haeIII/palI ACTGGGGCCA GTAACTGTCA AACTATGGAT TTGATACCTA CATTGACAGT TGACCCCGGT fokī sau96I nlaIV asuI tru9I maeIII SUBSTITUTE SHEET (RULE 26) 3901 3841

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62/70

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CCAAATACTG
                                                                                                                                                                                                                                                                                     GGTTTATGAC
                            mbol/ndell[dam-
                                                                                                                                                                            GGTTTGTTTG
                                                                                                                                                                                                                                                                           AGCGCAGATA
                                                                                                                                                                                     CCAAACAAAC
                                                                                               AGGAAAAAA
                                                                                                                                                                                                                                                                                     TCGCCTCTAT
                                                                                      TCCTTTTTT
                                                                                                                                                                                                                                                                  hhaI/cfoI
                                              dpnII[dam-
                                                                            bstYI/xhoII
                                     dpnI[dam+]
                                                        alwI[dam-]
                                                                                                                                                                                                                                                        hinPI
                   sau3AI
         mbol/ndell[dam-]
                                                                                      CTTCTTGAGA
                                                                                               GAAGAACTCT
                                                                                                                                                                                                                                                                            GCTTCAGCAG
                                                                                                                                                                                                                                                                                     CGAAGTCGTC
                                                                                                                                                                            TACCAGCGGT
                                                                                                                                                                                     ATGGTCGCCA
                                                                                                                                                                  nspBII
                                                                   mboll[dam-]
                                                                                                                                                                                                                                                                  eco57I
                            dpnII[dam-
                                      bstYI/xhoII
                   dpnI[dam+]
                                                alwI[dam-]
                                                         mbol/ndeII[dam-]
sau3AI
                                                                                                                                                                           AAACCACCGC
TTTGGTGGCG
                                                                                      CGTAGAAAG ATCAAAGGAT
                                                                                                TAGTTTCCTA
                                                                                                                                                                   acil
                                                                                                                                                                                                                                                                            AAGGTAACTG
                                                                                                                                                                                                                                                                                      TTCCATTGAC
                                                                                                                                                                                                                                                          bsrI
                                                                                                                                                                                                                                                                  maeIII
                                                                            dpnII[dam-]
                                                                   dpnI[dam+]
                                               sau3AI
                                                                                                GCATCTTTTC
                                                                                                                                                                                                                                                                                      AGAAAAAGGC
                                                                                                                                                                             GCAAACAAAA
                                                                                                                                                                                                                                                                             TCTTTTCCG
                                                                                                                                                                                       CGTTTGTTTT
                                                                                                                                                                                                                                                                                      TCGATGGTTG
                                                                                                GCAGTCTGGG
                                                                                                                                                                                                                                                                             AGCTACCAAC
                                                                                       CGTCAGACCC
                                                                                                                                                                             TCTGCTGCTT
                                                                                                                                                                                       AGACGACGAA
                                                                                                                                                           fnu4HI
                                                                                                                                                                                                                   mbol/ndeII[dam-]
                                                                                                                                                                    bbvI
                                                                                                                                                                                                                                                                    aluI
                                                                                                                              fnuDII/mvnI
                                                                    hgaI
                                                                                                                                                                                                                                       dpnII[dam-
                                                                                                                                                                                                                             dpnI[dam+]
                                                                                                                                                                                                                                                alwI[dam-]
                                                                                                                                                                    hhaI/cfoI
                                                                                                                                               bsh1236I
                                                                                       TTCCACTGAG
                                                                                                 AAGGTGACTC
                                                                                                                                                                                                                                                                                       GGCCTAGTTC
                                                                                                                                                                             CTGCGCGTAA
                                                                                                                                                                                      GACGCGCATT
                                                                                                                                                                                                                                                                             CCGGATCAAG
                                                                             ddeI
                                                                                                                                                                                                          sau3AI
                                                                                                                                       bstuI
              FIG. 16Y
                                                                                                                                                          hinPI
                                                                                                                    thal
                                                                                                                                                                                                                                                                   hpall
                                                                                                                                                                                                                                                          Idsm
                                                                                       4091
                                                                                                                                                                              4151
                                                                                                                                                                                                                                                                             4211
                                                                                                                         SUBSTITUTE SHEET (RULE 26)
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acil na cccccracar accrcccrcr sr gccccara rccacaga	scrfi ncil mspl hpall dsaV plel caull hinfl rA CCGGGTTGGA CTCAAGACGA	hgiAI/aspHI bsp1286 bsiHKAI bsiHKAI apaLI/snoI alw441/snoI alw441/snoI cACGTC ACAGCCCAGC CACGTG TGTCGGGTCG AACCTCGCTT	hinPI hhaI/cfoI aeII GCGCC ACGCTTCCCG AAGGGAGAA GCGCG TGCGAAGGGC TTCCCTCTT
scfi CTCTGTAGCA GAGACATCGT	TCGTGTCTTA	hgiAI/a bsp1286 bsiHKAI bmyI apaLI/s alw441/ GTTCGTGCAC	hinPI hhaI/c haeII AGAAAGCGCC TCTTTCGCGG
TCAAGAA	TGGCGATAAG	TGAACGGGGG	GTGAGCATTG
haell hael TTAGGCCACC ACT AATCCGGTGG TGA	fnu4HI alwNI bbvI bsrI fnu4HI III bbvI bsrI TACCAGTGG CTGCTGCCAG	nspBII fnu4HI bbvI mcrI nPI aciI aI/cfoI GCA GCGGTCGGGC	scfi TACCTACAGC ATGGATGTCG
bsli GTAGCCGTAG CATCGGCATC	fnu4HI alwNi bbvI bsrI fnu4HI eIII bbvI bsrI rraccagrgg crgcrgccag	nspBII fnu4HI bbvI mc hinPI aci hhaI/cfoI ATAAGGCGCA GCG TATTCCGCGT CGC	ddeI CGAACTGAGA GCTTGACTCT
rmal mael TCCTTCTAGT AGGAAGATCA	mae GCTAATCCTG T	mspI hpaII bsaWI maeIII TAGTTACCGG	CGACCTACAC
4281	4351	SUBSTITUTE SHEET (RULE 26	4491

FIG. I 6Z-

ecoRII mvaI ecoRII SCLFI dsaV mvaI

bstNI

aluI mnlI hhaI/cfoI hinPI

GAGCGCACGA

apyI[dcm+]

bsaJI

GGGAGCTTCC

CCCTCGAAGG CTCGCGTGCT CGGAACAGGA GCCTTGTCCT

CGCCGTCCCA

GCGCCAGGGT

TATCCGGTAA

GGCGGACAGG

4561

acil

CCCCCTGTCC

SCLFI

ATAGGCCATT

HHI

fun4

bslI

hpall

Idsm

٠

aciI

bsaWI

TCCCCCTTTG

AGGGGGAAAC

sfaNI taqI

TCGATTTTTG AGCTAAAAAC hgaI CTGAACTCGC GACTTGAGCG drdI CGCCACCTCT GCGGTGGAGA mnll ACAGCCCAAA TGTCGGGTTT

ACTACGAGCA

TGATGCTCGT

haeIII/palI fnu4HI acil

fnuDII/mvnI bslI thaI

bstuI

FIG. I 6Z-2

bsh1236I

GCAACGCGGC CGTTGCGCCG AAAAACGCCA TTTTTGCGGT GAGCCTATGG CTCGGATACC CAGGGGGGCG GICCCCCCGC

aciI

nlaIV

4701

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apyI[dcm+]

bstNI

dsaV

AAATATCAGG

CGGACCATAG

GCCTGGTATC

TTTATAGTCC

P.

65/70

GGGACTAAGA CCCTGATTCT hinfI tfil ACCGCCTCTC TGGCGGAGAG GACGCAATAG CCGAACGACC GGCTTGCTGG CTGCGTTATC mcrI mnlI aciI fnu4HI CAATACGCAA GTTATGCGTT TACAAGAAAG CTCGCCGCAG GAGCGGCGTC Inqq ATGTTCTTTC fnu4HI acil sapl hhal/cfol aflIII earI/ksp632I nspHI bsrBI nspI CTTCTCGCGG GAAGAGCGCC aciI hinPI CGACTATGGC haeII AAAACGAGTG GCTGATACCG TTTTGCTCAC nlaIII FIG. 1 6Z-3 Ilodm haeIII/palI aluI acil CGAGGAAGCG CTTTGAGTGA GAAACTCACT AAACGACCGG TTTGCTGGCC haeI mnlI haeIII/palI apyl[dcm+] TCAGTCACTC GAAAAATGCC AAGGACCGGA AGTCAGTGAG TICCIGGCCT acil GTATTACCGC CATAATGGCG bs1 haeI ecoRII bstNI SCLFI mvaI dsaV hinfI nlaIV pleI hhaI/cfoI GAGCGCAGCG CTCGCGTCGC CTTTTACGG GTGGATAACC CACCTATTGG fnu4HI **PpvI** hinPI 4871 4811 4741 SUBSTITUTE SHEET (RULE 26)

CACTTTATGC

GGTCCGAAAT

TAATCCGTGG

ATTAGGCACC

CCAGGCTTTA

66/70

acil I apyl[dcm+] bsaJI CTGGAAAGCG GACCTTTCGC ecoRII bstNI SCLFI dsaV mval bsrI GGTTTCCCGA (CCAAAGGGGCT ( hgiCI nlaIV banI TGGCACGACA ACCGTGCTGT mnlI maeIII nspBII aseI/asnI/vspI /asnI/vspI DVUII aluI TTAATCCAGC AATTAGGTCG I6n. mseI tru91 mseI aseI/ bsh1236I haeIII/palI CCCGCGCGTT GGCCGATTCA GGGCGCGCAA CCGGCTAAGT hinfI tfiI hhaI/cfoI fnuDII/mvnI eaeI cfrI hinPI fnuDII/mvnI **bsh1236I** hhaI/cfoI bstUI hinPI thaI bstUI thaI bslI 4931 SUBSTITUTE SHEET (RULE 26)

GGCAGTGAGC GCAACGCAAT TAATGTGAGT TACCTCACTC
CCGTCACTCG CGTTGCGTTA ATTACACTCA ATGGAGTGAG FIG. 162-4

ATTACACTCA ATGGAGTGAG

4991

ATGACCATGA TACTGGTACT GGAAACAGCT aluI ATTTCACACA TAAAGTGTGT GCGGATAACA aciI bsrBI GGAATTGTGA CCTTAACACT TATGTTGTGT ATACAACACA TTCCGGCTCG hpall ldsm

5061

nlalll

aseI/asnI/vspI

A

AATGCTTAAT

asp700 TTACGAATTA

5131

xmnI

tru9I

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mseI

FIG. I 62-5

5141 >length: WO 95/14930 PCT/US94/13329

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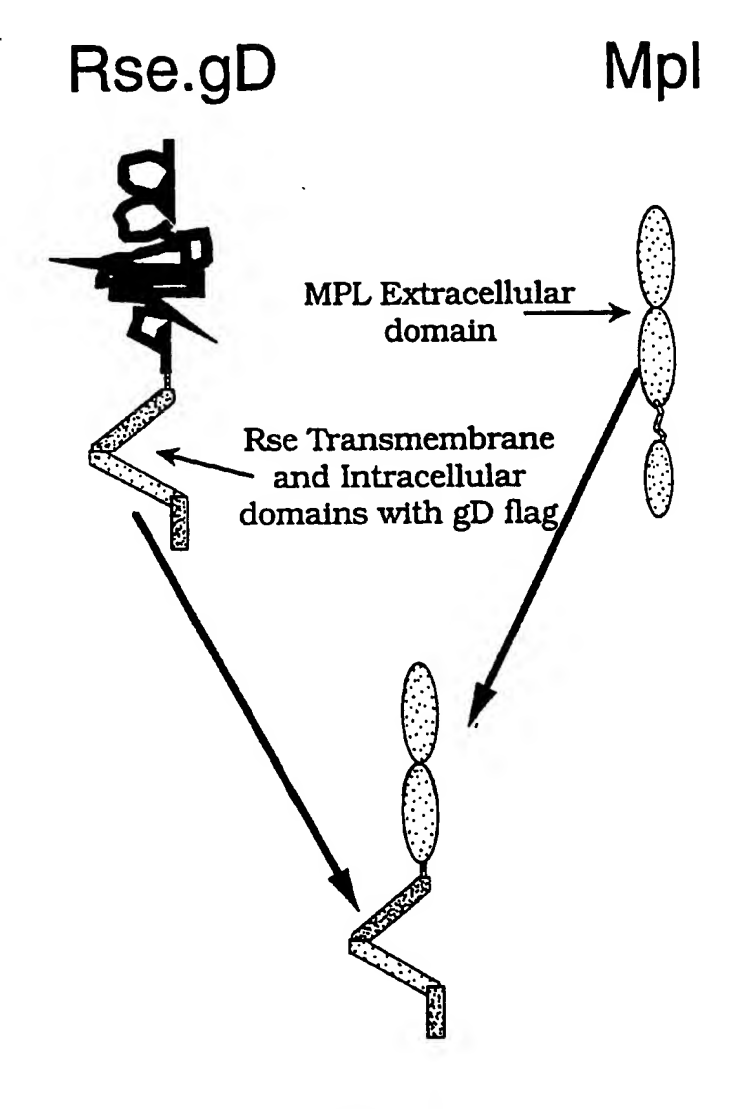
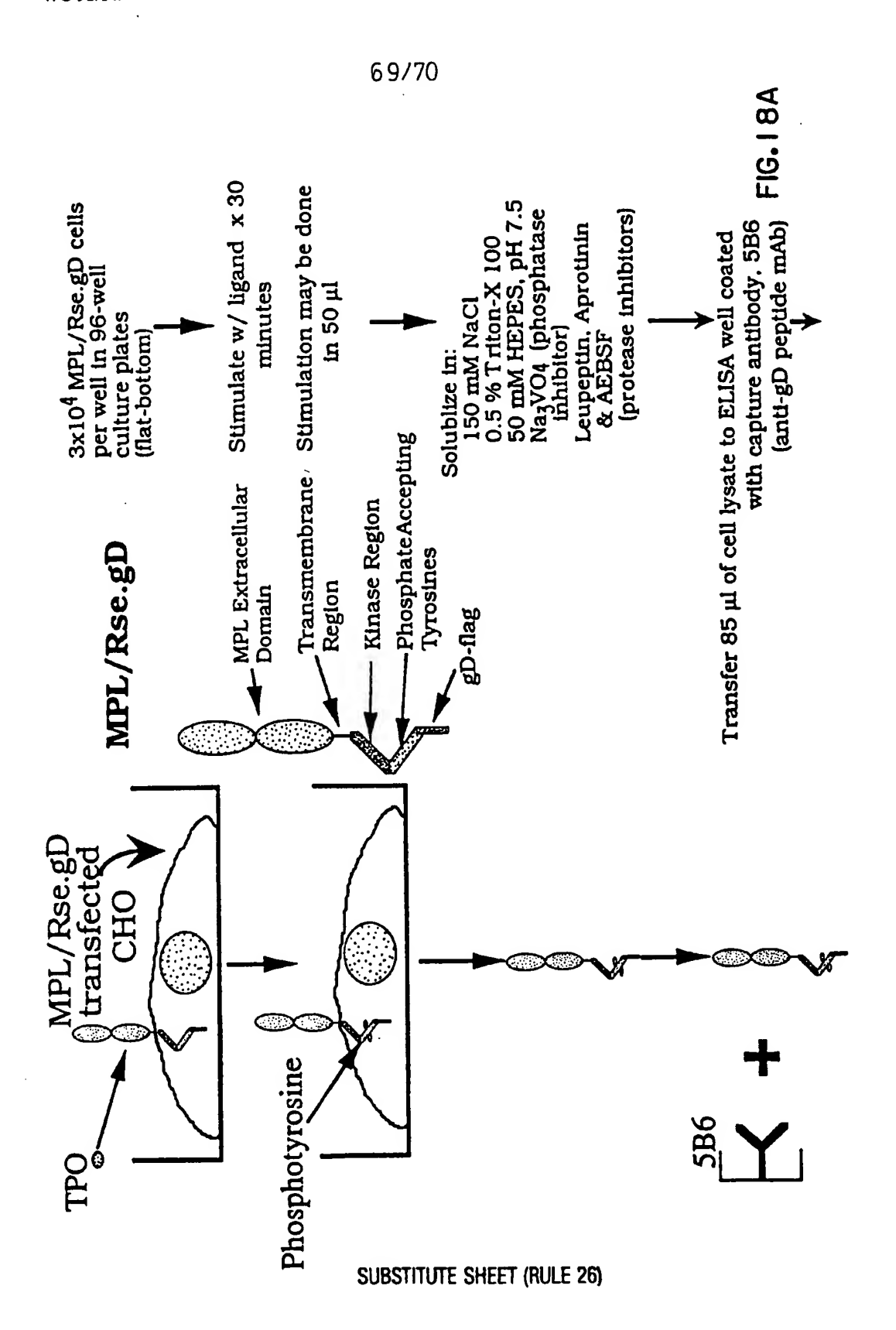
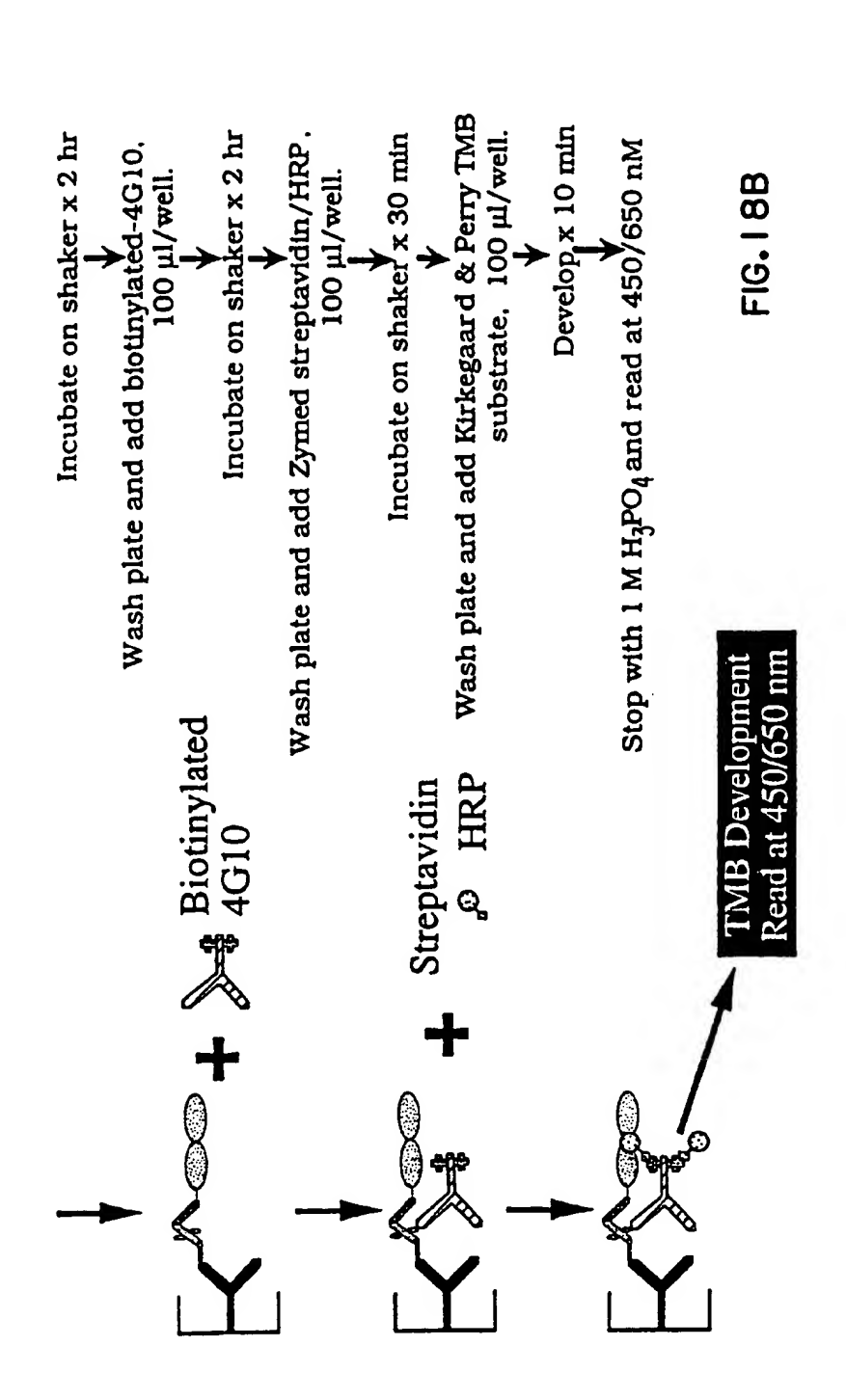


FIG. 17



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## INTERNATIONAL SEARCH REPORT

Internation Application No PCT/US 94/13329

	IFICATION OF SUBJECT MATTER G01N33/573 G01N33/566 G01N33, C07K14/705	/543 G01N33/532 G01N	33/577
According to	o International Patent Classification (IPC) or to both national cla	snification and IPC	
· · · · · · · · · · · · · · · · · · ·	SEARCHED		
Minimum d IPC 6	ocumentation searched (classification system followed by classification sy	cation symbols)	
	tion searched other than minimum documentation to the extent the		earched
C. DOCUM	MENTS CONSIDERED TO BE RELEVANT		Relevant to claim No.
Category *	Citation of document, with indication, where appropriate, of th	e relevant passages	KGEARIT M CIRTIL MO.
X,P	JOURNAL OF BIOLOGICAL CHEMISTRY vol.269, no.14, 1 April 1994, W DC USA pages 10720 - 10728 M.R. MARK ET AL. 'RSE, a novel receptor-type tyrosine kinase w homology to Ax1/Ufo, is express levels in the brain.' cited in the application see the whole document see the whole document	ASHINGTON with	1-43
V Fu	rther documents are listed in the continuation of box C.	X Patent family members are listed	in annex.
'A' docur consi 'E' earlier filing 'L' docur which citati 'O' docur other 'P' docur later  Date of the	ment defining the general state of the art which is not idered to be of particular relevance or document but published on or after the international grate ment which may throw doubts on priority claim(s) or this cited to establish the publication date of another ion or other special reason (as specified) ment referring to an oral disclosure, use, exhibition or means ment published prior to the international filing date but than the priority date claimed  1. April 1995	"T" later document published after the in or priority date and not in conflict verted to understand the principle or invention  "X" document of particular relevance; the cannot be considered novel or cannot involve an inventive step when the cannot be considered to involve an document of particular relevance; the cannot be considered to involve an document is combined with one or ments, such combination being obvious the art.  "&" document member of the same pate.  Date of mailing of the international.  2.7. 04. 95	iternational filing date with the application but theory underlying the se claimed invention of be considered to document is taken alone inventive step when the more other such document to a person skilled int family
	mailing address of the ISA	Authorized officer	
	European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl. Fax: (+31-70) 340-3016	Van Bohemen, C	

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
	ARCHIVES OF BIOCHEMISTRY AND BIOPHYSICS, vol.285, no.2, 1 March 1991, NEW YORK NY USA pages 197 - 204 V.P. KNUTSON ET AL. 'Comparison of insulin receptor tyrosine phosphorylation under in vitro and in situ conditions: assessment of specific protein tyrosine phosphorylation without the use of 32P-phosphate labeled substrates.' cited in the application see figure 1	1-28, 33-43
Y	DIABETES, vol.42, 1 June 1993, WASHINGTON DC USA pages 883 - 890 H.H. KLEIN ET AL. W 'A microtiter well assay system to measure insulin activation of insulin receptor kinase in intact human mononuclear cells.' cited in the application see page 883, abstract see page 884, column 2, line 9 - line 44	1-28, 33-43
	EP,A,O 244 221 (GENENTECH INCORPORATED) 4 November 1987 cited in the application see claim 1	1-28, 33-43
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Internatic Application No PCT/US 94/13329

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